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Gray et al.

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(54) **EXPANDABLE INTERVERTEBRAL IMPLANT**

USPC 606/246, 249, 279, 90, 105; 623/17.11, 623/17.15, 17.16

See application file for complete search history.

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(73) Assignee: **GLOBUS MEDICAL, INC.**, Audubon, PA (US)

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A61F 2/30 (2006.01)

A61F 2/46 (2006.01)

(52) **U.S. Cl.**

CPC **A61F 2/447** (2013.01); **A61F 2/442** (2013.01); **A61F 2/4455** (2013.01); (Continued)

(58) **Field of Classification Search**

CPC A61F 2/4455; A61F 2/442; A61F 2/447; A61F 2/4684; A61F 2/4611; A61F 2002/30062; A61F 2002/3008; A61F 2002/30181; A61F 2002/30266; A61F 2002/30387; A61F 2002/30405; A61F 2002/30448; A61F 2002/30476; A61F 2002/30556; A61F 2002/30579; A61F 2002/30607; A61F 2002/30616; A61F 2002/30677; A61F 2002/30787; A61F 2002/4475; A61F 2310/00023; A61F 2310/00029; A61F 2310/00359

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Primary Examiner — Pedro Philogene

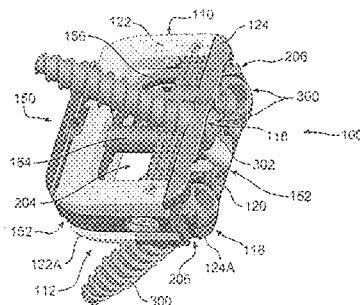
Assistant Examiner — David Comstock

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ABSTRACT

An implant for therapeutically separating bones of a joint has two endplates each having an opening through the endplate, and at least one ramped surface on a side opposite a bone engaging side. A frame is slideably connected to the endplates to enable the endplates to move relative to each other at an angle with respect to the longitudinal axis of the implant, in sliding connection with the frame. An actuator screw is rotatably connected to the frame. A carriage forms an open area aligned with the openings in the endplates. The openings in the endplates pass through the carriage to form an unimpeded passage from bone to bone of the joint. The carriage has ramps which mate with the ramped surfaces of the endplates, wherein when the carriage is moved by rotation of the actuator screw, the endplates move closer or farther apart.

15 Claims, 17 Drawing Sheets



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 (2013.01); *A61F 2002/3008* (2013.01); *A61F*
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 (2013.01); *A61F 2230/0082* (2013.01); *A61F*
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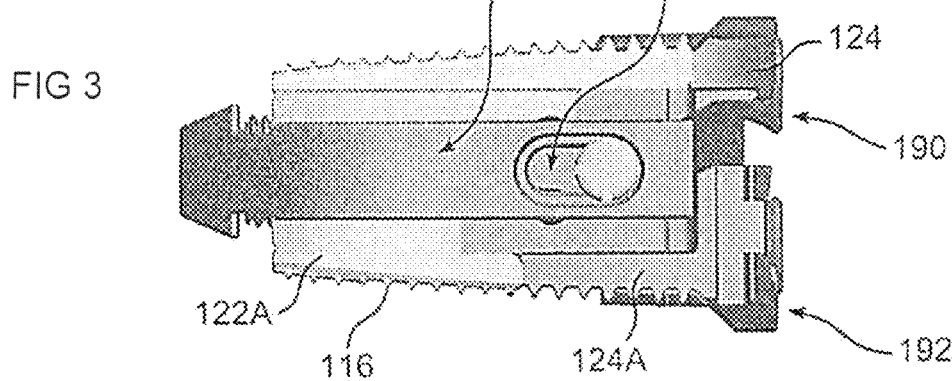
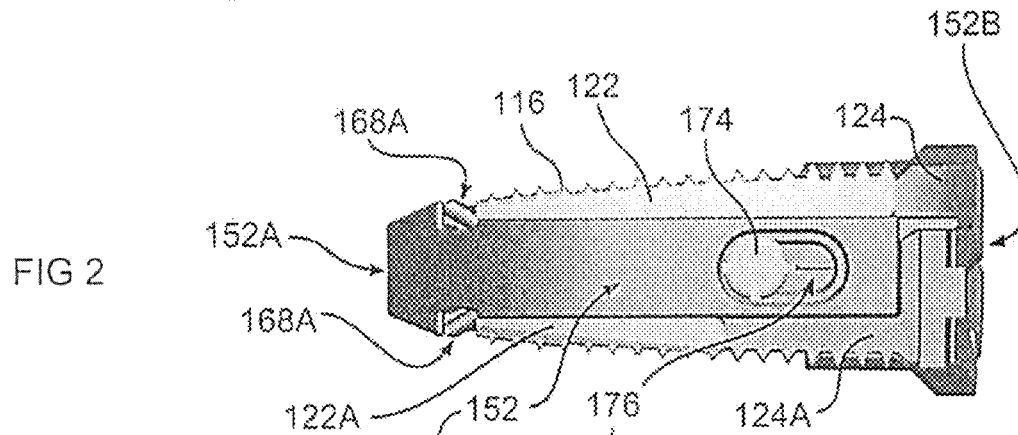
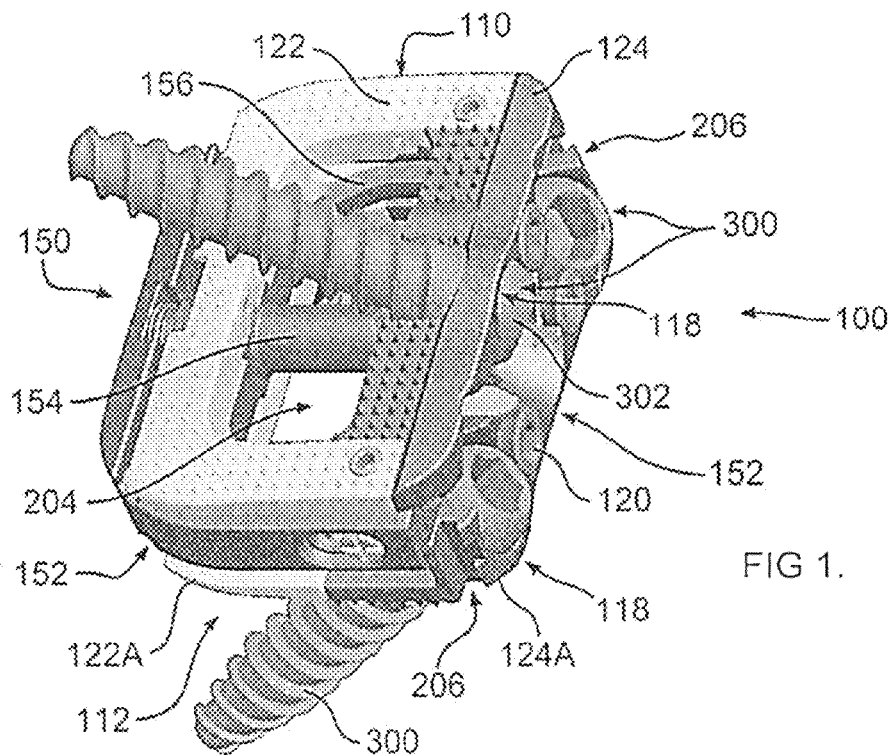
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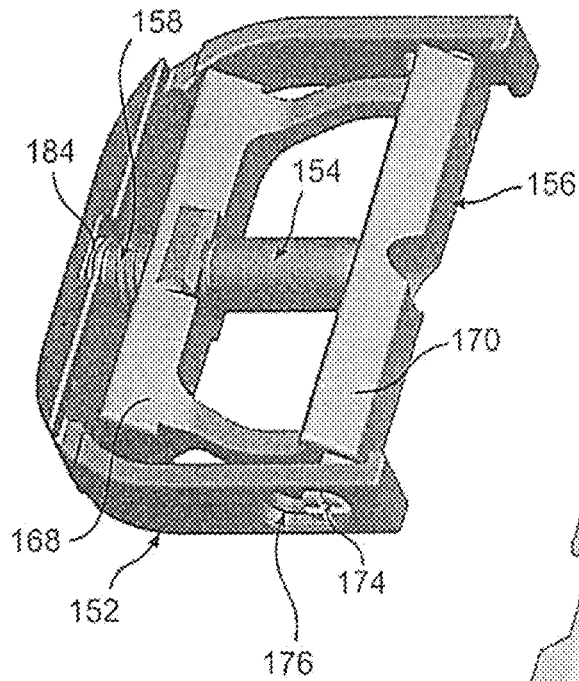


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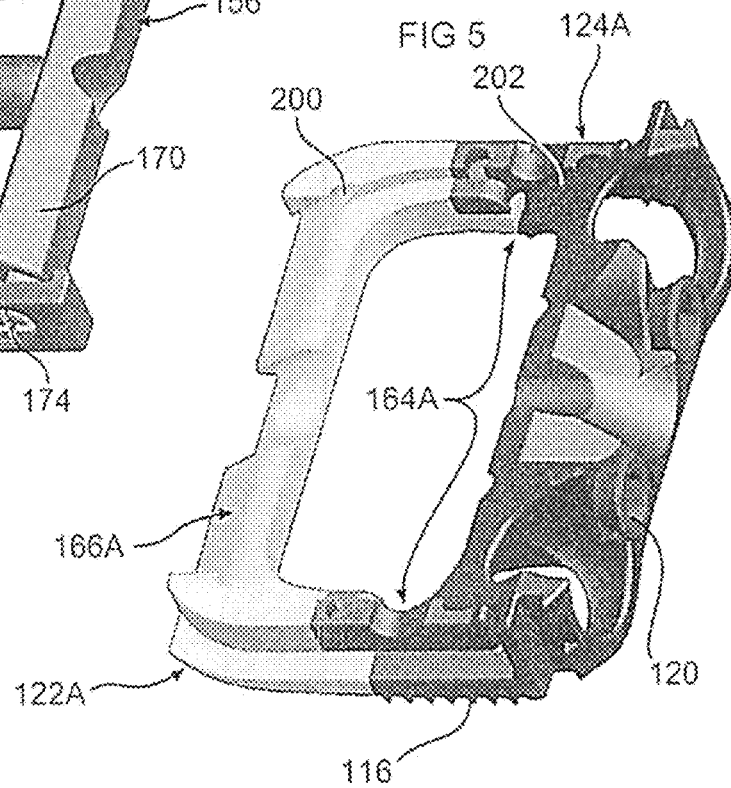


FIG 5

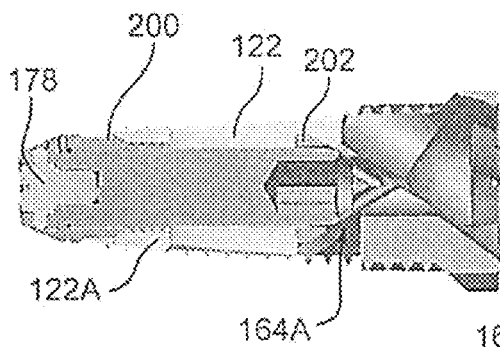
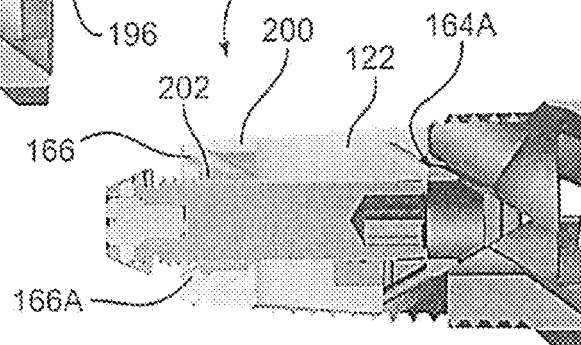
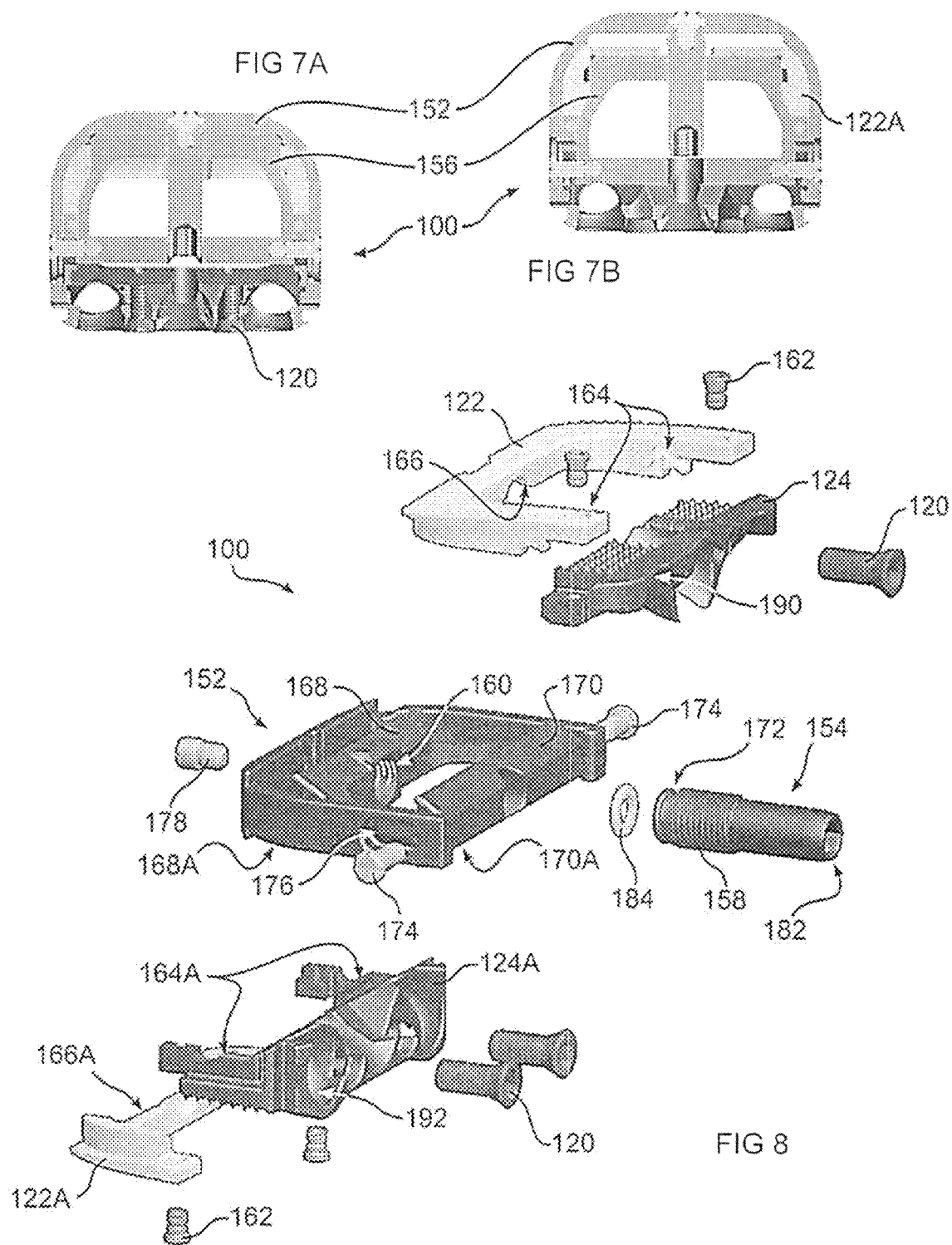
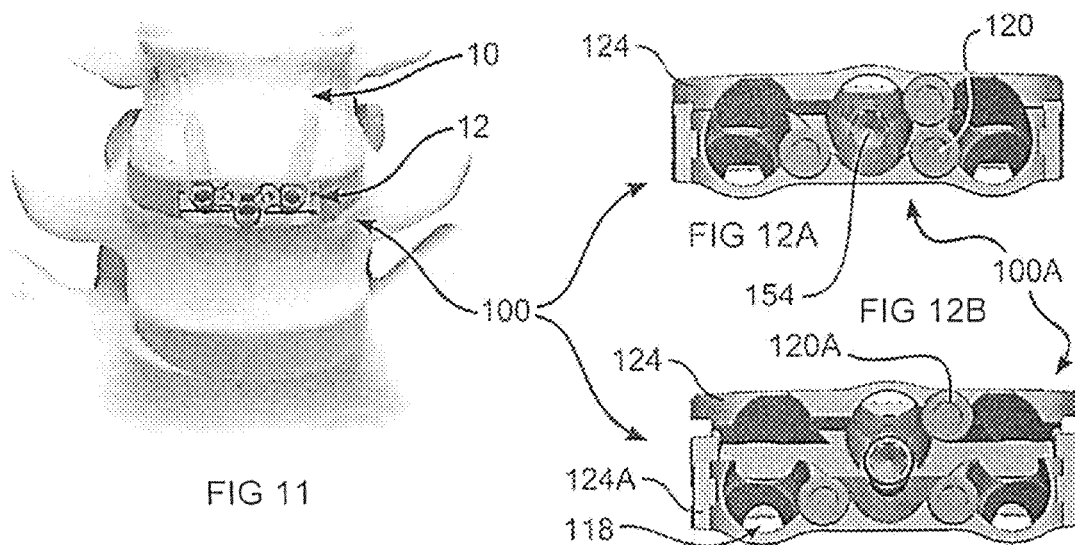
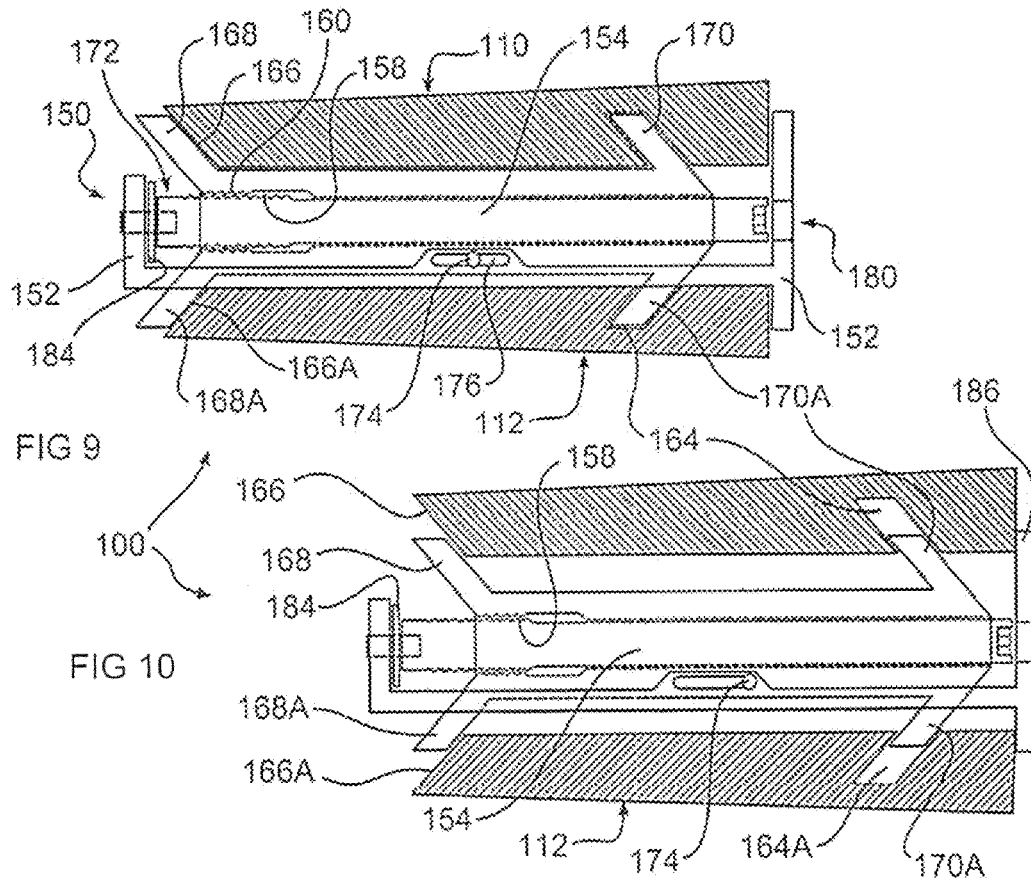


FIG 6A

FIG 6B







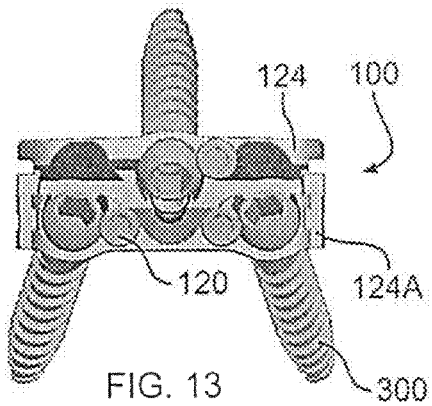


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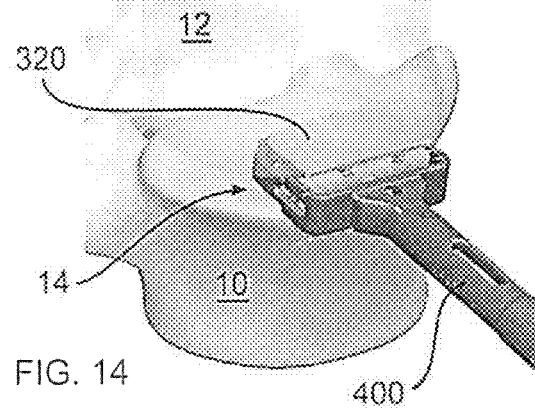


FIG. 14

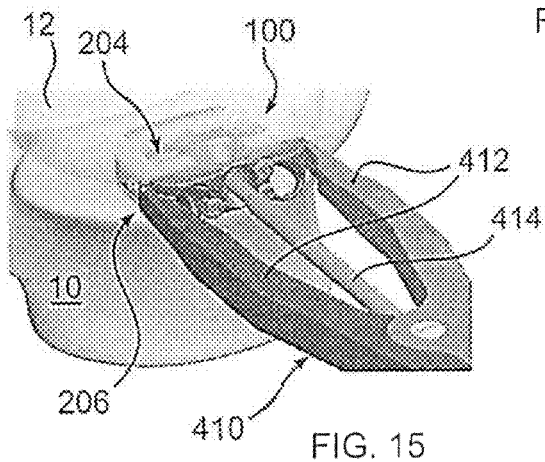


FIG. 15

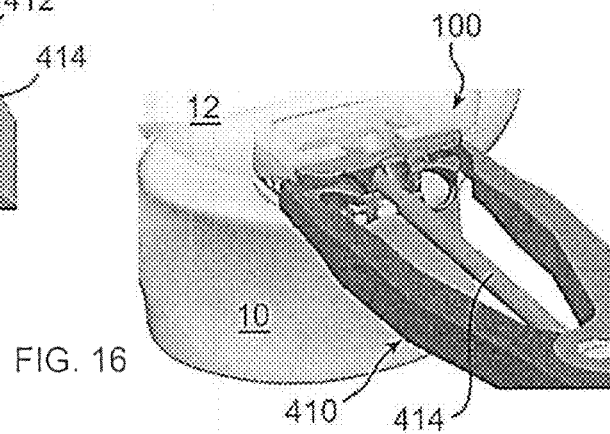


FIG. 16

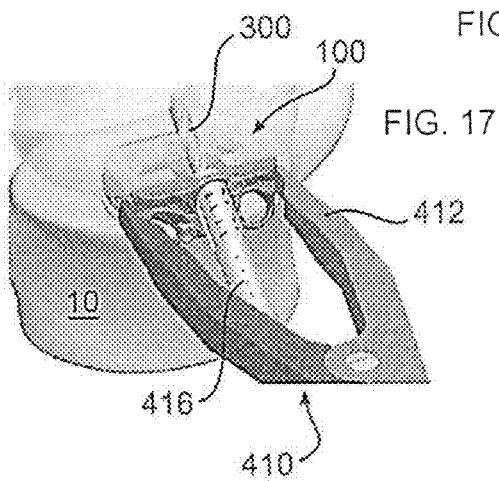


FIG. 17

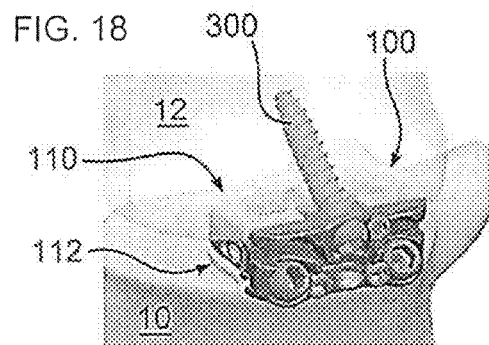
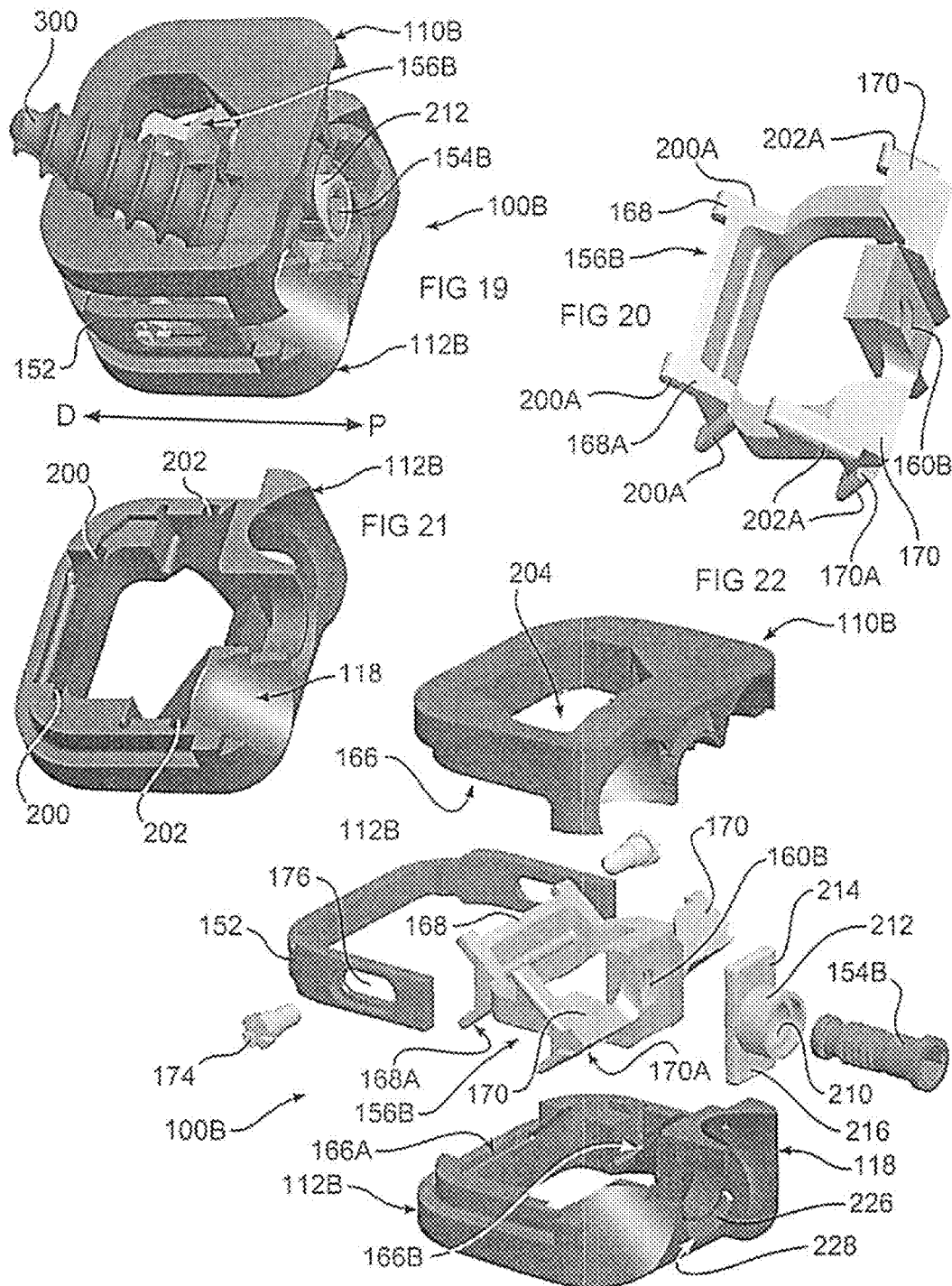


FIG. 18



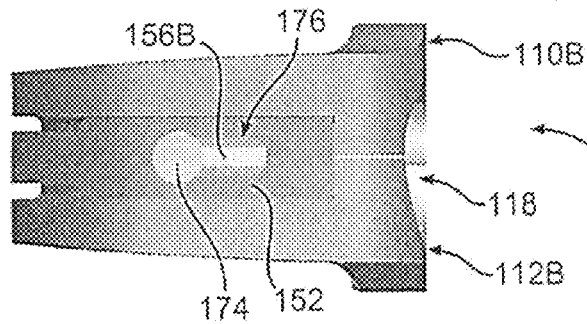


FIG 23

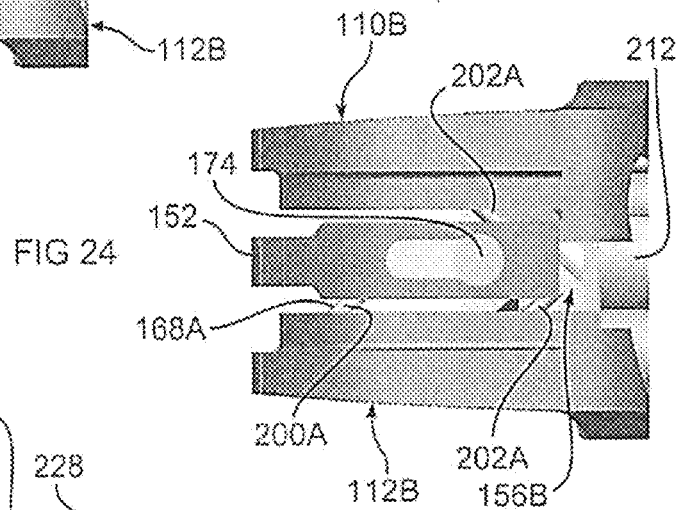


FIG 24

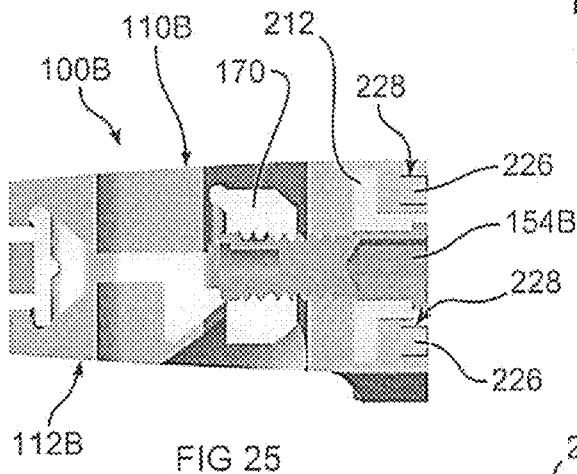


FIG 25

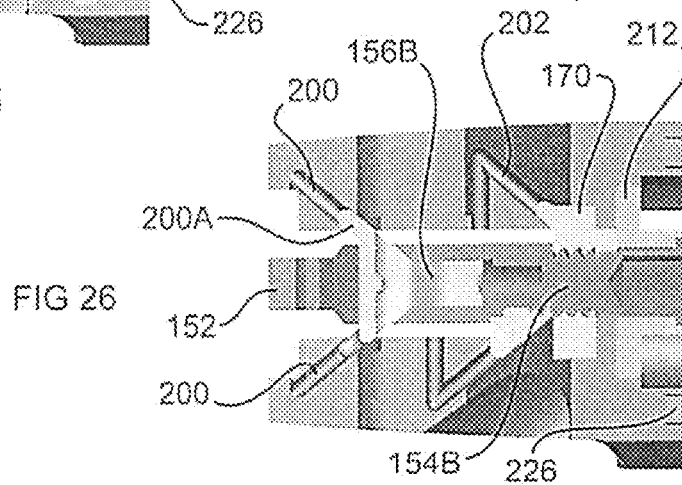
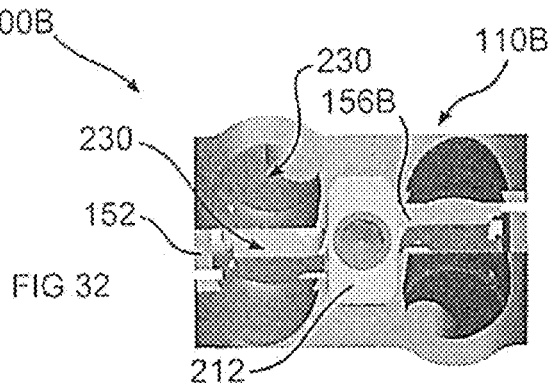
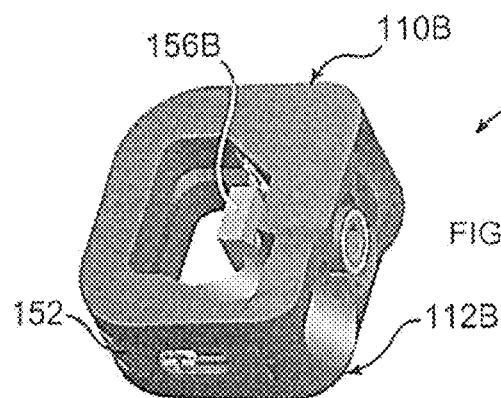
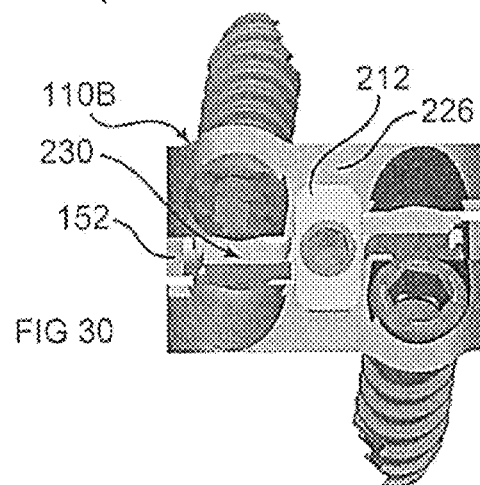
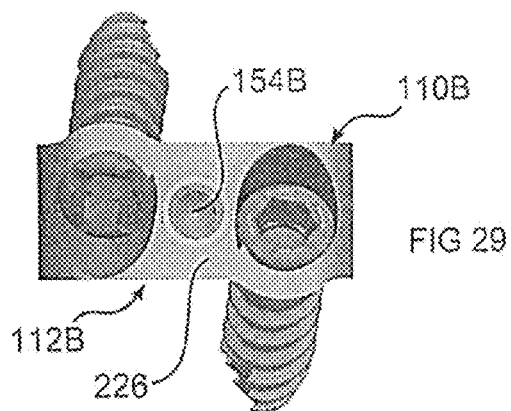
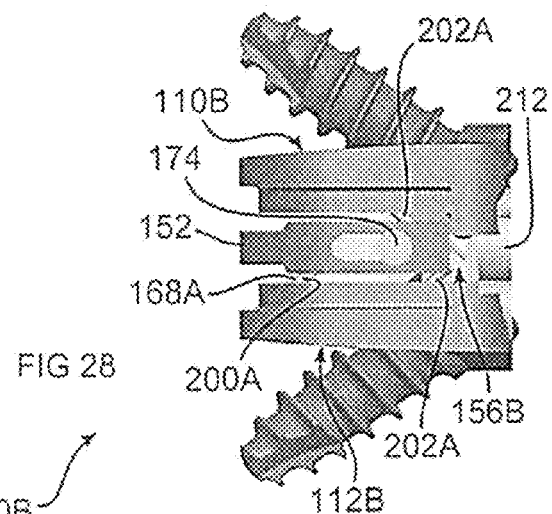
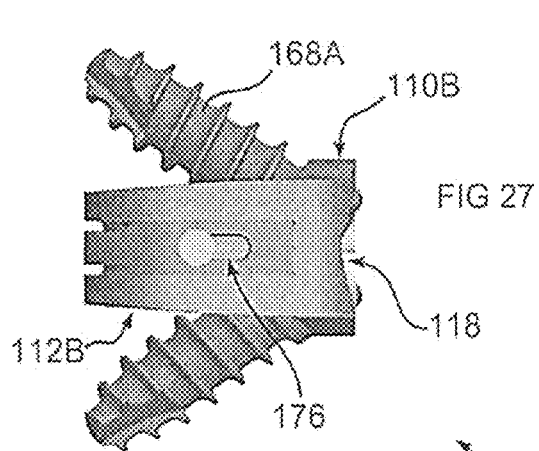
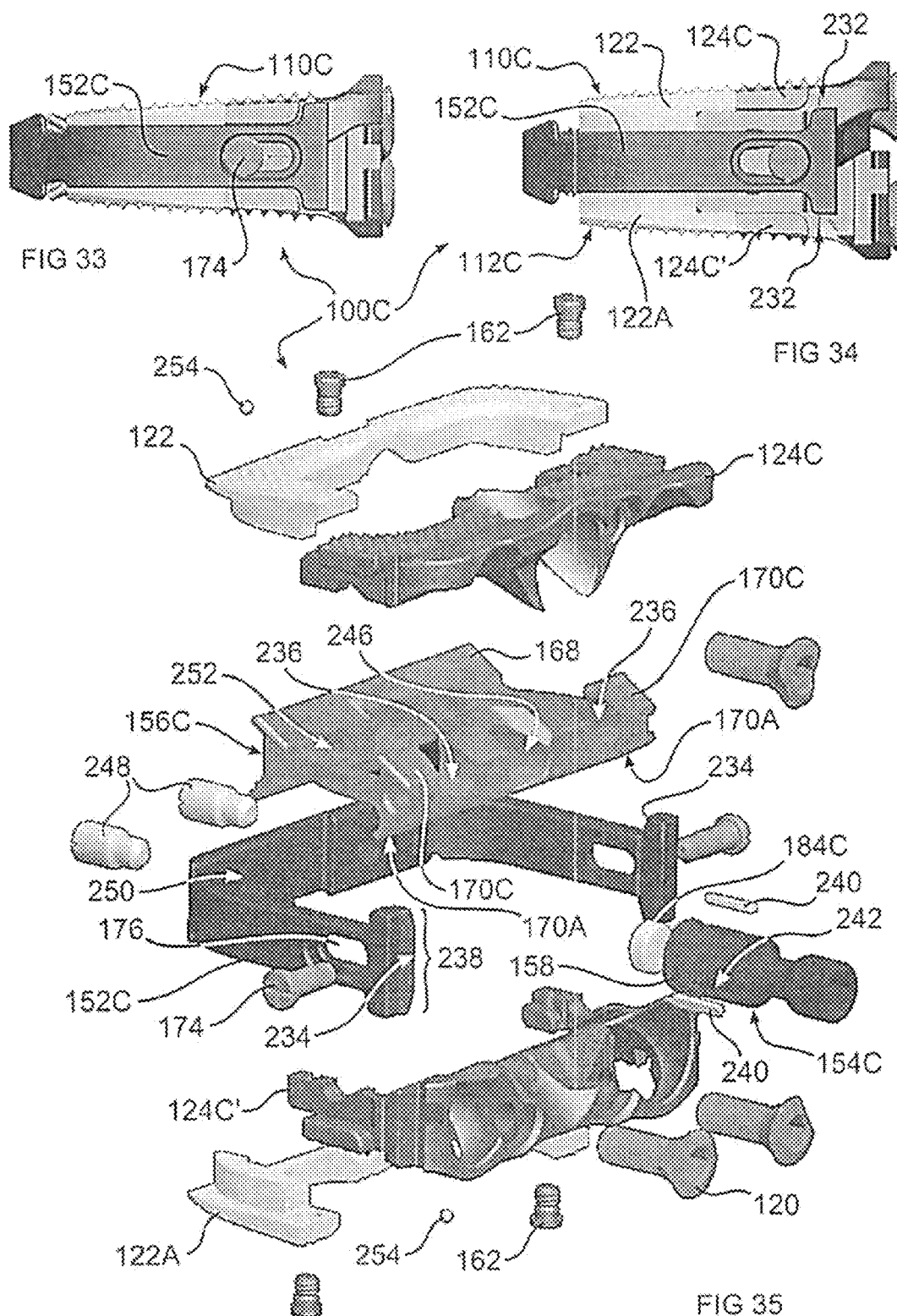
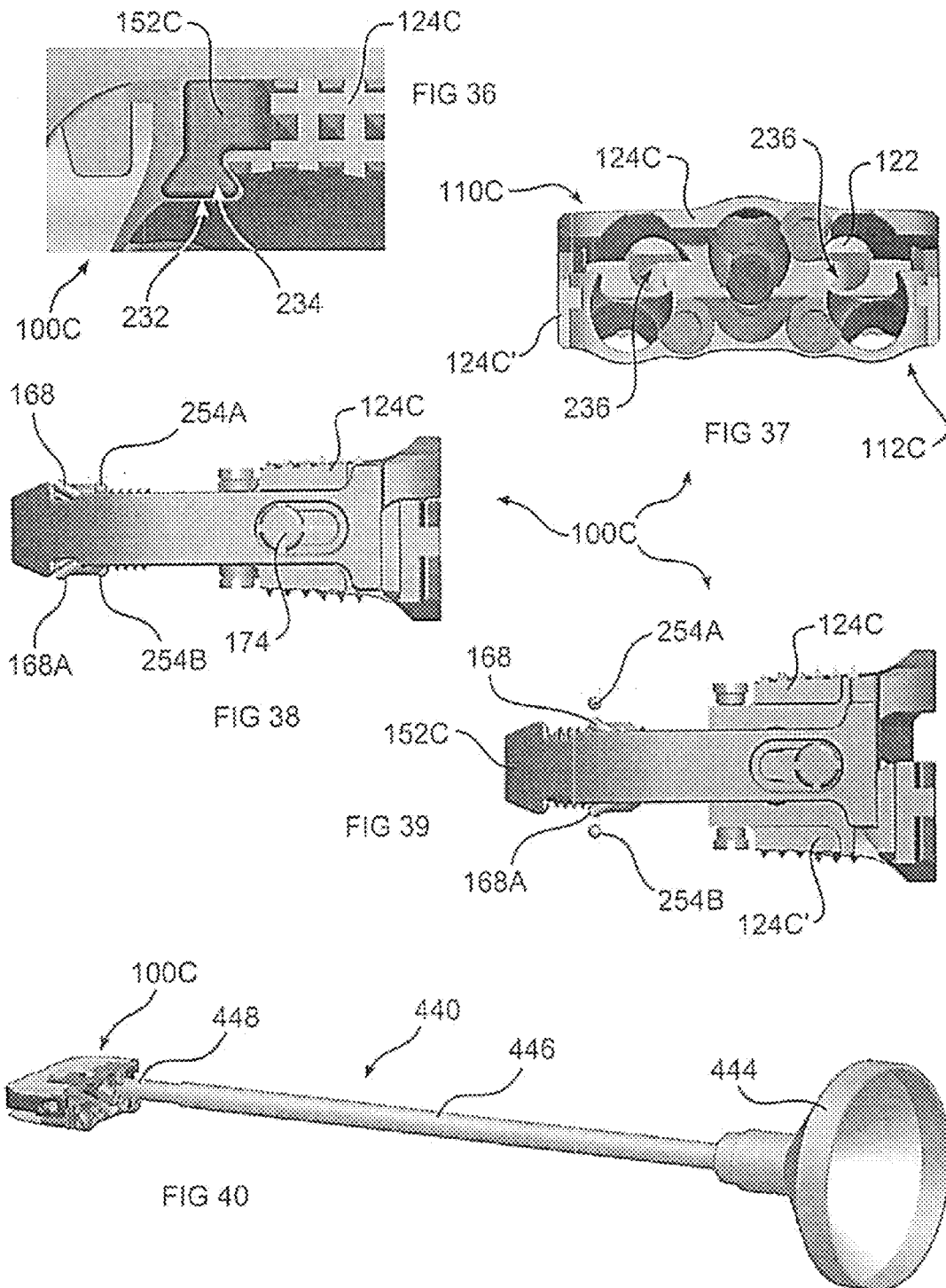


FIG 26







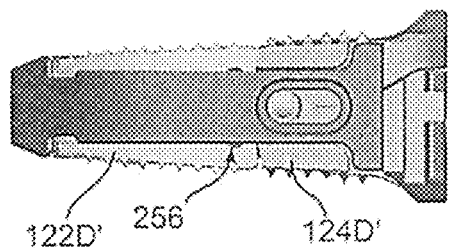


FIG 41

FIG 42

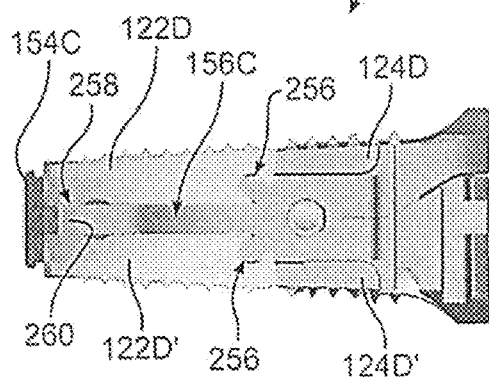
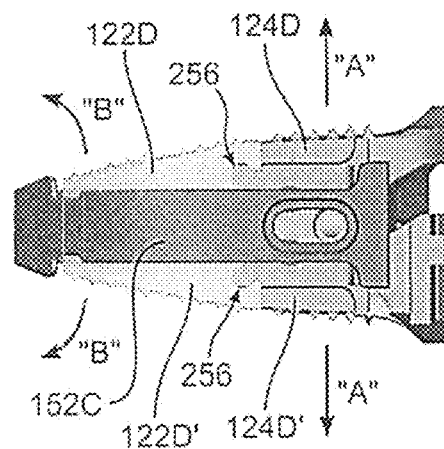


FIG 43

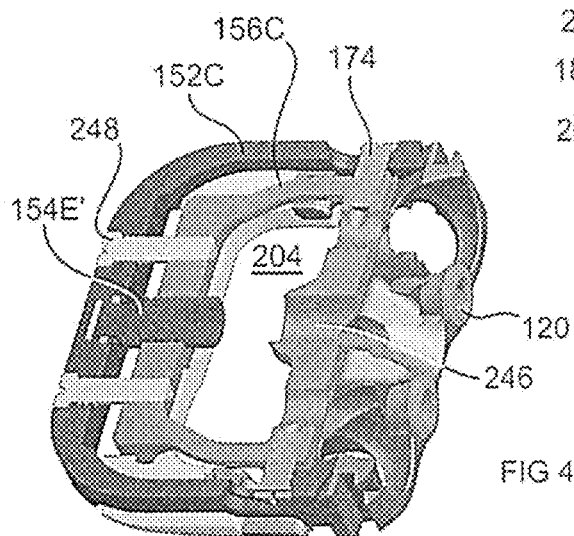


FIG 45

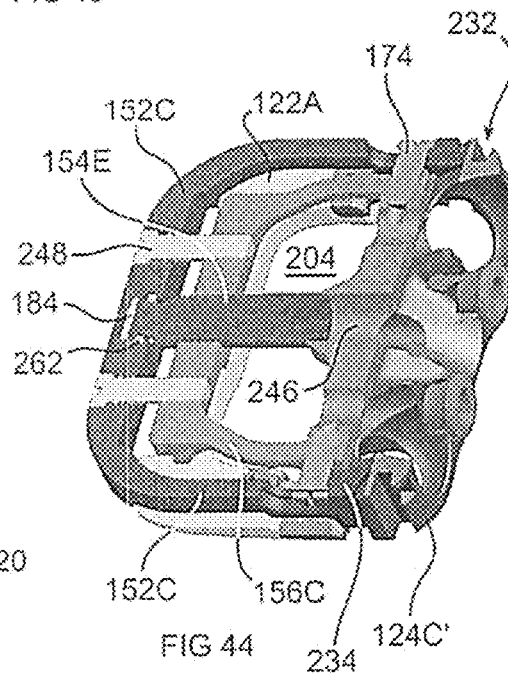
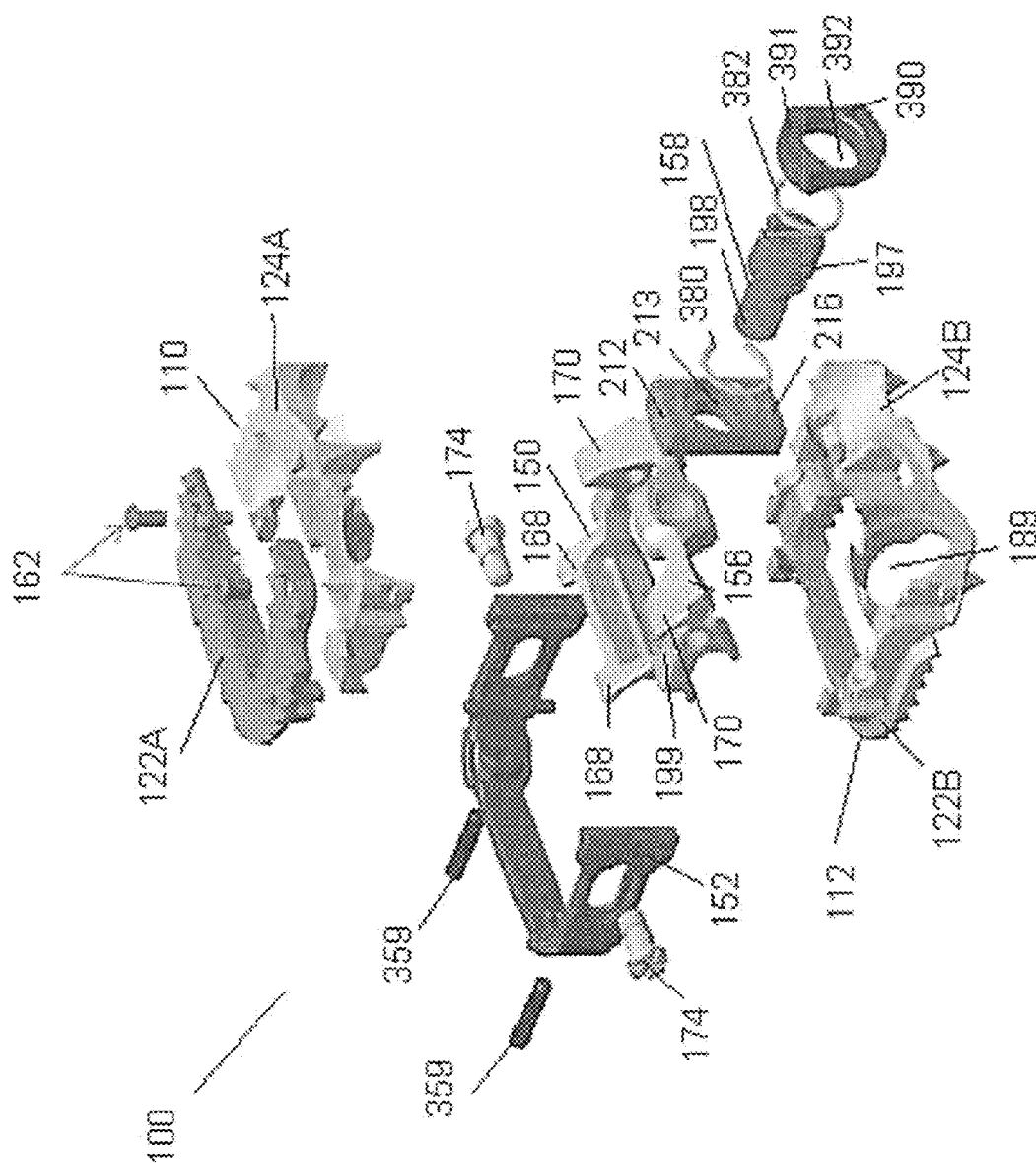


FIG 44



64
65

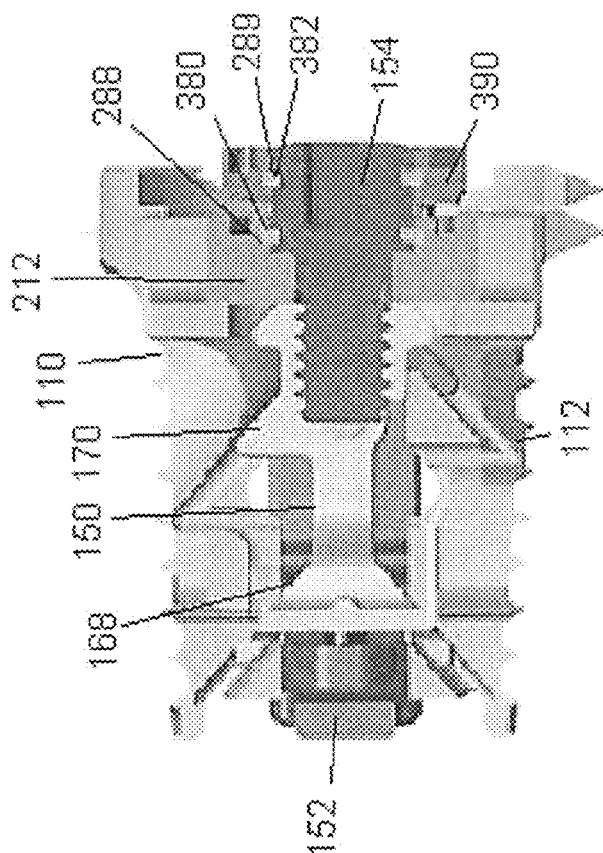


FIG. 47B

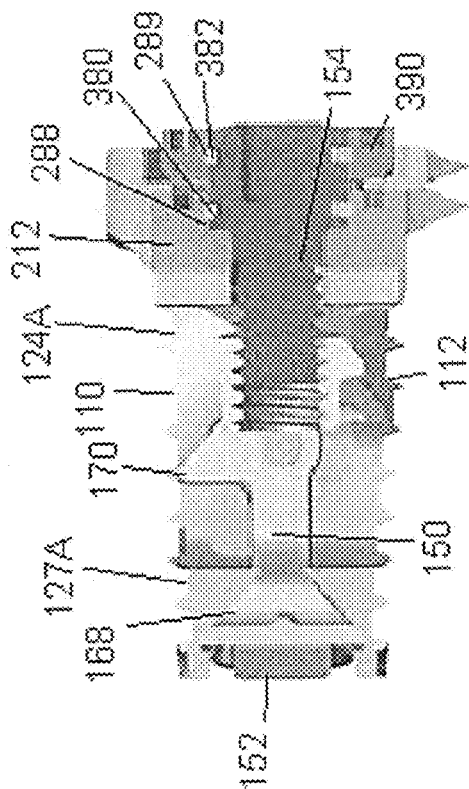


FIG. 47A

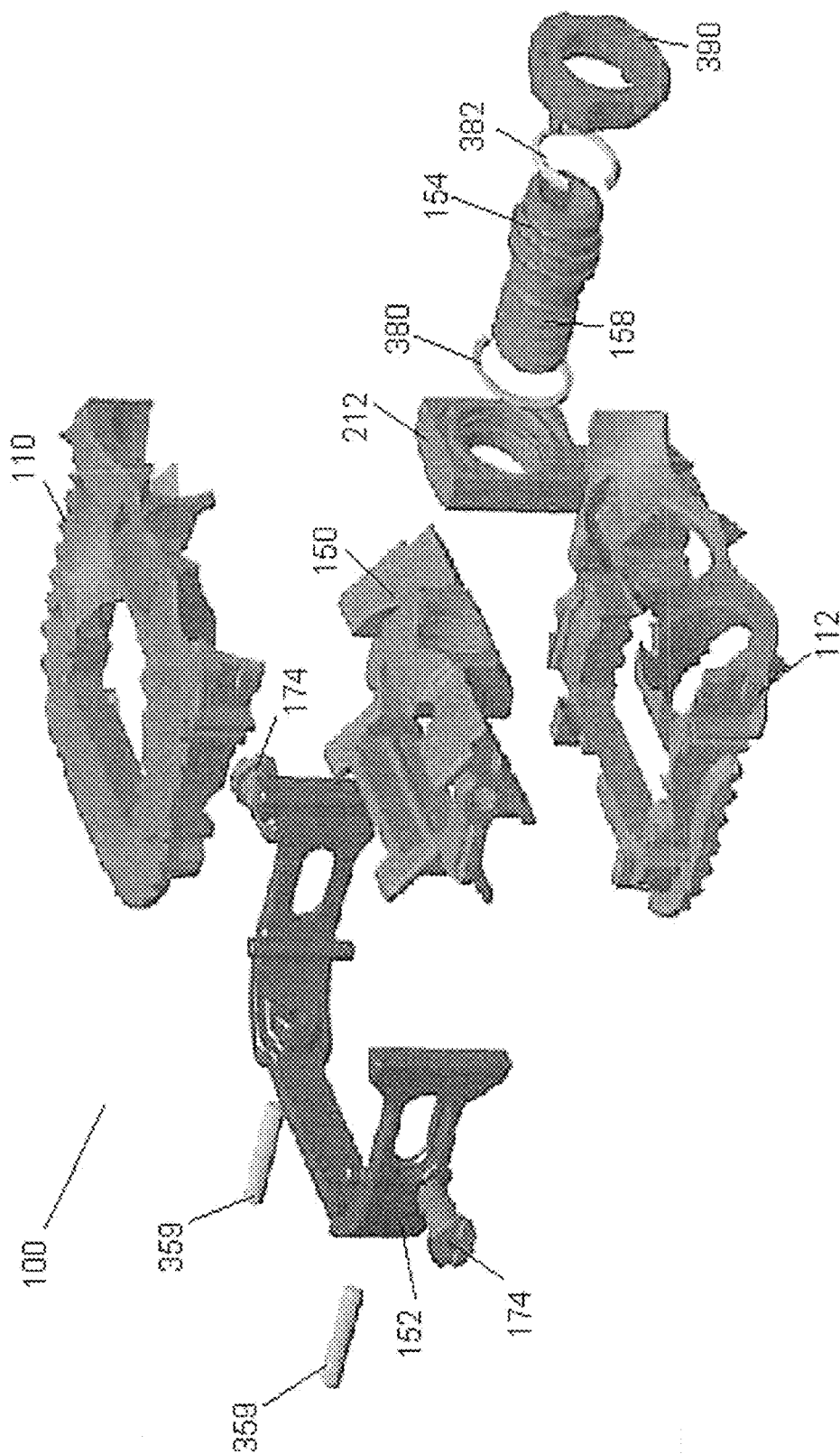


FIG. 48

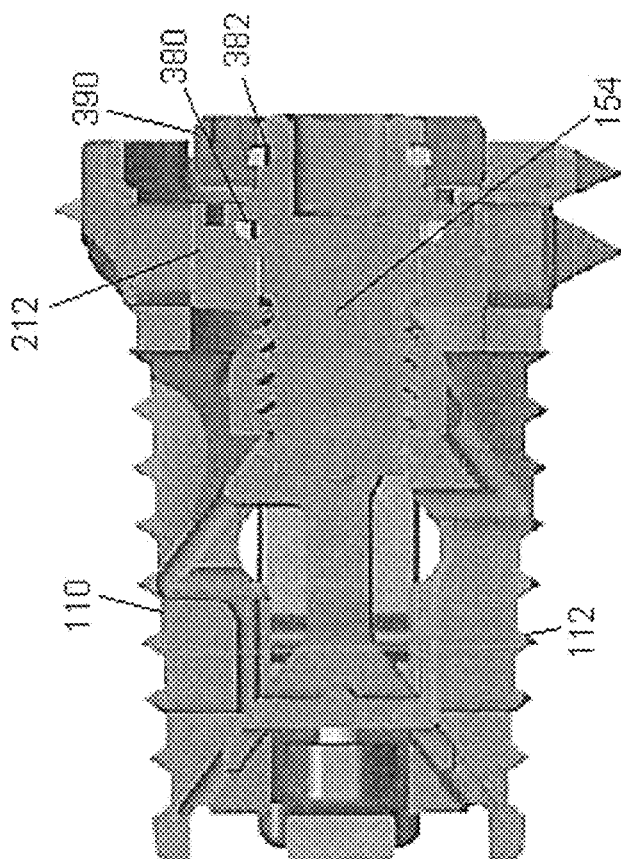


FIG. 49A

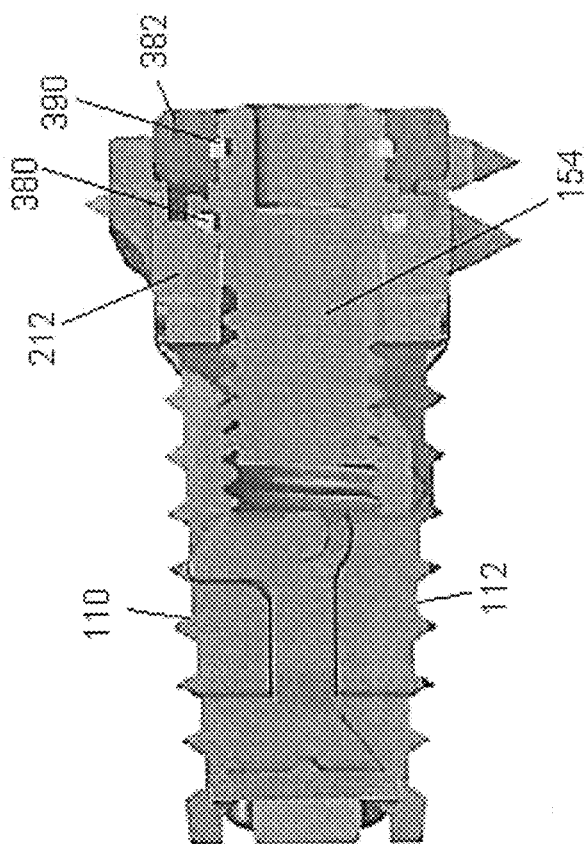


FIG. 49B

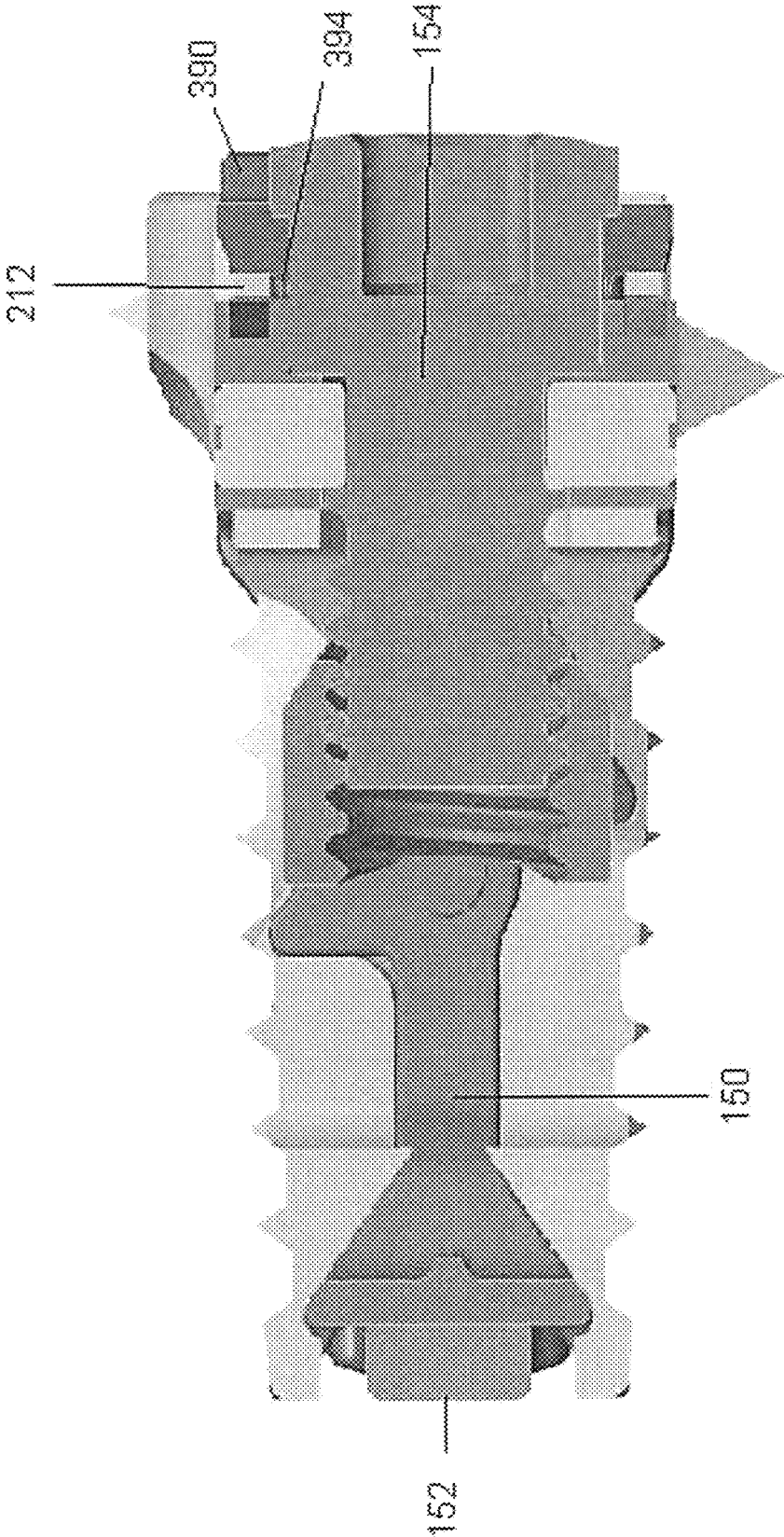


FIG. 50A

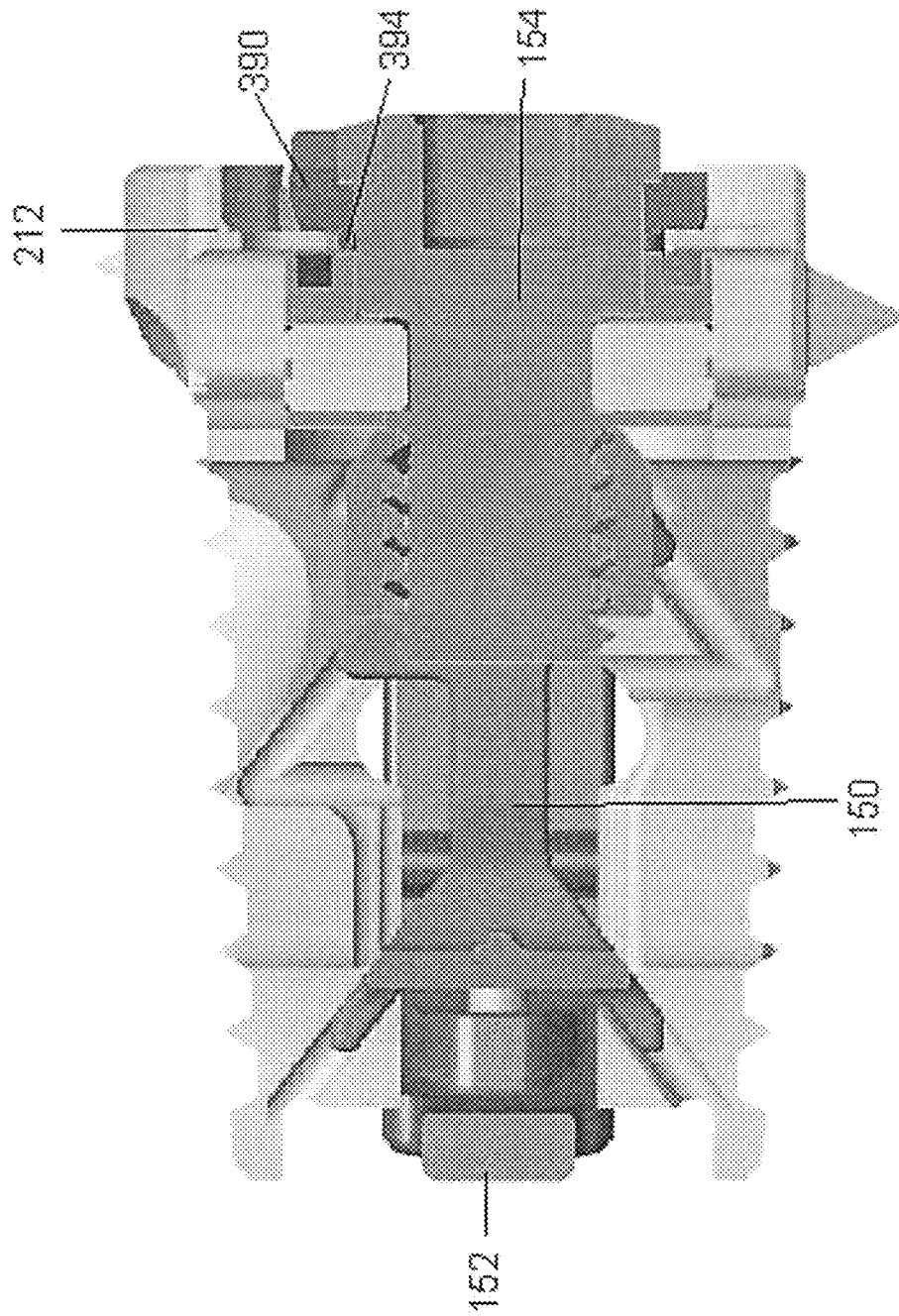


FIG. 50B

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**EXPANDABLE INTERVERTEBRAL
IMPLANT****CROSS-REFERENCE TO RELATED
APPLICATIONS**

This application is a continuation-in-part of U.S. patent application Ser. No. 13/836,687, entitled "Expandable Intervertebral Implant," filed on Mar. 15, 2013, for which the disclosure is incorporated herein by reference in its entirety.

FIELD OF THE INVENTION

This invention relates to stabilizing adjacent vertebrae of the spine by inserting an intervertebral implant, and more particularly an intervertebral implant that is adjustable in height.

BACKGROUND OF THE INVENTION

Bones and bony structures are susceptible to a variety of weaknesses that can affect their ability to provide support and structure. Weaknesses in bony structures have numerous potential causes, including degenerative diseases, tumors, fractures, and dislocations. Advances in medicine and engineering have provided doctors with a plurality of devices and techniques for alleviating or curing these weaknesses.

In some cases, the spinal column requires additional support in order to address such weaknesses. One technique for providing support is to insert a spacer between adjacent vertebrae.

SUMMARY OF THE INVENTION

In accordance with the disclosure, an implant for therapeutically separating bones of a joint, the implant defining a longitudinal axis extending between distal and proximal ends, the implant comprises a first endplate configured to engage a first bone of the joint, and having an opening through the endplate, and at least one ramped surface on a side opposite a bone engaging side; a second endplate configured to engage a second bone of the joint, and having an opening through the endplate, and at least one ramped surface on a side opposite a bone engaging side; a frame slideably connected to the first and second endplates to enable the first and second endplates to move relative to each other at an angle with respect to the longitudinal axis, in sliding connection with the frame; an actuator screw rotatably connected to the frame; and a carriage (a) forming an open area aligned with the openings in the first and second endplates and defining thereby a proximal carriage side and a distal carriage side with respect to the longitudinal axis, (b) threadably connected to the actuator screw, whereby rotation of the actuator screw moves the carriage with respect to the frame and the first and second endplates, the actuator screw not crossing between the proximal carriage side and the distal carriage side; and (c) including a plurality of ramps each mateable with at least one of the at least one ramped surfaces of the first and second endplates, wherein when the carriage is moved by rotation of the actuator screw, at least one of the at least one ramped surface of the first endplate and at least one of the at least one ramped surface of the second endplate each slide along at least one of the plurality of ramps of the carriage to cause the endplates to move relative to each other in sliding connection with the frame.

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In various embodiments thereof, the first and second endplates are confined by the frame to move relative to each other only along an axis substantially transverse to the longitudinal axis; at least one of the first and second endplates includes at least one aperture through which a fastener may pass to secure the implant to bone of the joint; the implant further includes a blocking mechanism configured to prevent backing out of a fastener passed through at least one of the first and second endplates and into body tissue; the blocking mechanism includes a blocking member slideably retained within a channel between an unblocking position and a blocking position in which a portion of the blocking member overlaps a portion of the faster; at least one of the first and second endplates includes one or more projections configured to engage bone of the joint when the implant is positioned between bones of the joint; at least one of the first and second endplates is composed of two interconnected portions of dissimilar materials; one of the dissimilar materials is metallic and includes at least one aperture through which a fastener may be passed to attach the implant to a bone of the joint; one dissimilar material is polymeric, and another dissimilar material is metallic; and, the implant further includes a polymeric material configured to press against the actuator screw to reduce a potential for unintended rotation of the actuator screw.

In further embodiments thereof, when the actuator screw is rotated in a first direction, a height of the implant transverse to the longitudinal axis is increased, and when the actuator screw is rotated in a second direction, a height of the implant transverse to the longitudinal axis is decreased; the actuator screw is threadably connected to the carriage along a proximal side of the carriage; the frame extends from the proximal end of the implant to the distal end of the implant, and the actuator screw is connected to the frame and threadably connected to the carriage along a distal side of the carriage; the frame is disposed within the proximal end of the implant; the frame extends from the proximal end of the implant towards the distal end of the implant; and, the implant further includes at least one post extending through the frame and into the carriage, slideably received in one of the frame or the carriage, thereby configured to maintain an alignment of the carriage along the longitudinal axis.

In yet further embodiments thereof, the implant further includes a first passage formed in a proximal end of at least one of the first and second endplates, and a second passage formed in a proximal side of the carriage, the first and second passages aligned to admit introduction of a therapeutic matter into the open area of the carriage when the implant is implanted between bones of the joint; the frame connects to the first and second endplates with a dovetail connection; the implant further includes at least one radiopaque marker positioned in connection with at least one of the first and second endplates, whereby an extent of movement of the connected endplate can be determined using imaging by a relative alignment of the radiopaque marker and a radiopaque element of the implant which does not move together with the connected endplate; ends of the at least one of the plurality of ramps of the carriage slide within grooves in at least one of the first and second endplates.

In another embodiment thereof, the frame includes an actuator screw bearing, a first tab extending away from the bearing in a first direction, and a second tab extending away from the bearing in a direction opposite to the upper tab, the first and second tabs forming edges; and the first and second

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endplates including grooves sized and dimensioned to slidably receive the edges of the first and second tabs, respectively.

In accordance with another embodiment of the disclosure, an implant for therapeutically separating bones of a joint, the implant defining a longitudinal axis extending between distal and proximal ends, the implant comprises a first endplate configured to engage a first bone of the joint, and having an opening through the endplate transverse to the longitudinal axis, and at least one ramped surface on a side opposite a bone engaging side; a second endplate configured to engage a second bone of the joint, and having an opening through the endplate transverse to the longitudinal axis, and at least one ramped surface on a side opposite a bone engaging side;

a frame slideably connected to the first and second endplates to enable the first and second endplates to move relative to each other at an angle substantially transverse to the longitudinal axis, in sliding connection with the frame; an actuator screw rotatably connected to the frame; and a carriage (a) forming an open area aligned with the openings in the first and second endplates and defining thereby a proximal carriage side and a distal carriage side with respect to the longitudinal axis, (b) threadably connected to the actuator screw, whereby rotation of the actuator screw moves the carriage with respect to the frame and the first and second endplates, the actuator screw not crossing between the proximal carriage side and the distal carriage side; (c) including a plurality of ramps each mateable with at least one of the at least one ramped surfaces of the first and second endplates, wherein when the carriage is moved by rotation of the actuator screw, at least one of the at least one ramped surface of the first endplate and at least one of the at least one ramped surface of the second endplate each slide along at least one of the plurality of ramps of the carriage to cause the endplates to move relative to each other in sliding connection with the frame; and (d) at least one passage formed in a proximal side of the carriage in communication with at least one proximal passage in at least one of the first or second endplates, the communicating passages configured to admit introduction of a therapeutic matter into the open area of the carriage when the implant is implanted between bones of the joint.

In accordance with the disclosure, a method of therapeutically separating bones of a joint, comprises inserting an implant defining a longitudinal axis extending between distal and proximal ends between bones of the joint, the implant including—a first endplate configured to engage a first bone of the joint, and having an opening through the endplate, and at least one ramped surface on a side opposite a bone engaging side; a second endplate configured to engage a second bone of the joint, and having an opening through the endplate, and at least one ramped surface on a side opposite a bone engaging side; a frame slideably connected to the first and second endplates to enable the first and second endplates to move relative to each other at an angle with respect to the longitudinal axis, in sliding connection with the frame; an actuator screw rotatably connected to the frame; and a carriage (a) forming an open area aligned with the openings in the first and second endplates and defining thereby a proximal carriage side and a distal carriage side with respect to the longitudinal axis, (b) threadably connected to the actuator screw, whereby rotation of the actuator screw moves the carriage with respect to the frame and the first and second endplates, the actuator screw not crossing between the proximal carriage side and the distal carriage side; and (c) including a plurality of ramps

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each mateable with at least one of the at least one ramped surfaces of the first and second endplates, wherein when the carriage is moved by rotation of the actuator screw, at least one of the at least one ramped surface of the first endplate and at least one of the at least one ramped surface of the second endplate each slide along at least one of the plurality of ramps of the carriage to cause the endplates to move relative to each other in sliding connection with the frame; and rotating the actuator screw after the implant is inserted to move the first and second endplates relatively farther apart to separate bones of the joint.

BRIEF DESCRIPTION OF THE DRAWINGS

A more complete understanding of the present invention, and the attendant advantages and features thereof, will be more readily understood by reference to the following detailed description when considered in conjunction with the accompanying drawings, in which:

FIG. 1 depicts an implant of the disclosure, together with three mounted bone screws;

FIG. 2 depicts the implant of FIG. 1, in a compressed or reduced height configuration;

FIG. 3 depicts the implant of FIG. 1, in an expanded or increased height configuration;

FIG. 4 depicts a carriage and frame of the implant of FIG. 1;

FIG. 5 depicts an endplate of the implant of FIG. 1;

FIG. 6A depicts a sagittal cross-section of the implant of FIG. 2;

FIG. 6B depicts a sagittal cross-section of the implant of FIG. 3;

FIG. 7A depicts an transverse cross-section of the implant of FIG. 2;

FIG. 7B depicts an transverse cross-section of the implant of FIG. 3;

FIG. 8 depicts an exploded view of the implant of FIG. 1;

FIG. 9 depicts a diagrammatic view of aspects of an implant in accordance with the disclosure, in a reduced height configuration;

FIG. 10 depicts the implant of FIG. 9, in an expanded height configuration;

FIG. 11 depicts the implant of FIG. 1, implanted between adjacent vertebrae;

FIG. 12A depicts a front view of the implant of FIG. 1 having an alternative blocking configuration, in a reduced height configuration;

FIG. 12B depicts the implant of FIG. 12A in an expanded height configuration;

FIG. 13 depicts the implant of FIG. 12B, with bones screws inserted into the implant;

FIG. 14 depicts inserting a trial of the disclosure, the trial representing an implant of the disclosure, into the disc space, using a trialing tool of the disclosure;

FIG. 15 depicts an implantation and actuating tool of the disclosure inserting an implant of the disclosure into the disc space;

FIG. 16 depicts the implant and tool of FIG. 14, the tool having expanded the implant;

FIG. 17 depicts the implant and tool of FIG. 15, and a bone screw driver inserting a bone screw;

FIG. 18 depicts the implant of FIG. 13 secured between vertebrae;

FIG. 19 depicts an implant of the disclosure including a proximally driven carriage;

FIG. 20 depicts the carriage of the implant of FIG. 19;

FIG. 21 depicts a lower endplate of the implant of FIG. 19;

FIG. 22 depicts an exploded view of the implant of FIG. 19;

FIG. 23 depicts a reduced height configuration of the implant of FIG. 19;

FIG. 24 depicts an expanded height configuration of the implant of FIG. 23;

FIG. 25 depicts a cross section of the implant of FIG. 23;

FIG. 26 depicts a cross section of the implant of FIG. 24;

FIG. 27 depicts the implant of FIG. 23, with bone screws inserted into the implant;

FIG. 28 depicts the implant of FIG. 24, with bone screws inserted into the implant;

FIG. 29 depicts a front view of the implant of FIG. 27;

FIG. 30 depicts a front view of the implant of FIG. 28;

FIG. 31 depicts a perspective view of the implant of FIG. 19, without bone screws inserted;

FIG. 32 depicts a front view of the implant of FIG. 30, without bone screws inserted;

FIG. 33 depicts a side view of an alternative implant in accordance with the disclosure, in a reduced height configuration;

FIG. 34 depicts the implant of FIG. 33, in an expanded height configuration;

FIG. 35 depicts an exploded view of the implant of FIG. 33;

FIG. 36 depicts an enlarged cross section of a dovetail connection of the implant of FIG. 33;

FIG. 37 depicts a front view of the implant of FIG. 33, illustrating passages for bone graft material;

FIG. 38 depicts a simulating of radiographic imaging of an implant of the disclosure, illustrating radiographic markers, the implant in a reduced height configuration;

FIG. 39 depicts the implant of FIG. 38, the implant in an expanded height configuration;

FIG. 40 depicts a bone funnel of the disclosure, used in connection with an implant of the disclosure;

FIG. 41 depicts an alternative implant of the disclosure, including hinged endplates, in a reduced height configuration;

FIG. 42 depicts the implant of FIG. 41, in an expanded configuration;

FIG. 43 depicts the implant of FIG. 41, with a frame portion removed;

FIG. 44 depicts a cross section of an alternative implant of the disclosure, in perspective, having an elongate actuator screw;

FIG. 45 depicts the implant of FIG. 44, having a shortened actuator screw;

FIG. 46 depicts an exploded view of an alternative expandable implant including guide pins in accordance with embodiments of the present application;

FIGS. 47A and 47B depict unexpanded and expanded configurations of the expandable implant in FIG. 46;

FIG. 48 depicts an exploded view of an alternative expandable implant including endplates formed primarily of metal in accordance with embodiments of the present application;

FIGS. 49A and 49B depict unexpanded and expanded configurations of the expandable implant in FIG. 48; and

FIGS. 50A and 50B depict unexpanded and expanded configurations of an alternative expandable implant having an alternate means to capture the blocking mechanism in accordance with embodiments of the present application.

DETAILED DESCRIPTION OF THE INVENTION

As required, detailed embodiments are disclosed herein; however, it is to be understood that the disclosed embodiments are merely examples and that the systems and methods described below can be embodied in various forms. Therefore, specific structural and functional details disclosed herein are not to be interpreted as limiting, but merely as a basis for the claims and as a representative basis for teaching one skilled in the art to variously employ the present subject matter in virtually any appropriately detailed structure and function. Further, the terms and phrases used herein are not intended to be limiting, but rather, to provide an understandable description of the concepts.

The terms “a” or “an”, as used herein, are defined as one or more than one. The term plurality, as used herein, is defined as two or more than two. The term another, as used herein, is defined as at least a second or more. The terms “including” and “having,” as used herein, are defined as comprising (i.e., open language).

Implants of the disclosure allow continuous expansion and retraction within a range of expansion. Lordosis of certain embodiments of implants herein can be custom tailored to fit the anatomy of a specific patient. Additionally, implants of the disclosure enable distraction of vertebral bodies to a desired height, but can also be collapsed and repositioned, as therapeutically indicated for the patient.

With reference to FIGS. 1-3, implant or implant 100 is operative, when positioned between adjacent bones of a joint, such as for example vertebrae 10, 12 (shown in FIG. 11), to stabilize a joint formed by adjacent vertebrae. Implant 100 has a collapsed state or height, illustrated in FIG. 2, and an expanded state or height, illustrated in FIG. 3. Implants 100 of the disclosure may be inset into the intervertebral disc space at a collapsed height, and then expand axially (superior/inferior) to restore height loss in the disc space. The implant provides distraction as well as achieves optimal height restoration. When inserted in a collapsed state, implants 100 reduce impaction to tissue in the joint space during insertion, and form the least visually blocking or obstructing profile.

Implant 100 includes two separable endplates 110, 112. A surface 114 of an endplate 110, 112 can be provided with teeth or other projections 116 which can penetrate body tissue to reduce a likelihood of migration of implant 100 after implantation. Implant 100 is further secured with one or more bone screws 300, which pass through bone screw socket 118 within implant 100, and into body tissue of the patient. In the embodiment illustrated in FIGS. 1-3, three sockets 118 for three bone screws are provided, the bone screws 300 further retained in connection with implant 100 by blocking fasteners 120. Bone screw 300 can be a polyaxial screw, and sockets 118 correspondingly shaped, whereby bone screw 300 may be inserted into body tissue at an optimal angle with respect to implant 100, whereby optimal purchase may be obtained, or certain body tissue may be avoided.

Endplates 110, 112 are moveably connectable to an actuator 150 operable to change a relative relationship of endplates 110 and 112. Actuator 150 includes a frame 152 rotatably supporting an actuator screw 154, and a moveable carriage 156. As actuator screw 154 rotates within frame 152, carriage 156 slides within frame 152, driven by cooperation between threads 158 (FIG. 8) upon actuator screw 154, and mating threads 160 within carriage 156. In the embodiment of FIGS. 1-3, endplates 110 and 112 are formed

in two connected portions, including a portion 122, 122A which can be polymeric, and a portion 124, 124A, which can be metallic. The portions are joined in the embodiment shown by screws 162, although other methods of combining the two connected portions 122, 124 or 122A and 124A may be used, including a dovetail connection, or adhesive, possibly in combination with each other, or with endplate connector screws 162. Metallic portions 124, 124A can provide greater strength for portions of implant 100 which are under relatively greater stress, for example portions through which a fastener may pass to anchor implant 100 within the body. While portions 122, 122A, 124, 124A are described as polymeric or metallic, it should be understood that other materials may be used, and that the portions can be of dissimilar materials.

With reference to FIG. 2, it may be seen that implant 100 is in a compressed state, having a lower height relative to an expanded state, as shown in FIG. 3. A functioning of device 100 may be best understood with reference to FIGS. 9-10, which correlate with FIGS. 2-3, respectively, but which present a simplified view having certain elements eliminated or exaggerated, to ease understanding. Endplates 110 and 112 are provided with ramped channels 164, 164A, and an open ramp 166, 166A, sized to slidably receive ramps 168, 168A and 170, 170A disposed upon carriage 156. While two mating channels and ramps are illustrated for each endplate 110, 112, it should be understood that one, or more than two, sets of channels and or ramps may be provided. Further, channels 164, 164A may alternatively be formed as ramps. However, a channel can operate to enable a reduction of height, having an opposing ramp face, whereby rotation of actuator screw 154 in an opposite direction to expansion can drive endplates 110, 112 together, for example when pressure from body tissue is insufficient to collapse endplates 110, 112. Additionally, at least one channel can operate to foster the maintenance of a connection between carriage 156 and an endplate 110, 112.

Carriage 156 is supported by frame 152 by lateral engagement means, in this embodiment two support screws 174 engaged with carriage 156, and passable through respective channels 176 formed in frame 152. Distal end 172 of actuator screw 154 provides additional support for carriage 156. Actuator screw 154 is supported by a set screw 178, which passes through and is rotatably supported within frame 152.

An actuator access port 180 permits passage of a tool, for example a hex driver (not shown), into engagement with a proximal end 182 of actuator screw 154. As actuator screw 154 is turned, distal end 172 bears against a thrust washer 184, and an end portion of frame 152. As actuator screw 154, carriage 156 is driven along actuator screw by interaction of threads 158 and 160. As carriage 156 moves, endplates 110, 112 are urged to move along ramps 168, 168A and 170, 170A, moving relatively apart, and increasing a height of implant 100. Endplates 110, 112 are prevented from moving together with carriage 156 by abutting against an end portion 186 of frame 152. In a given orientation, one of endplate 110 and 112 is an upper endplate with respect to an orientation in a standing patient. However, implant 100 may, in some embodiments, be implantable in either of opposite orientations, and therefore designations of upper and lower are provided for ease of understanding, only. It should be understood that only one of endplate 110, 112 may be moveable with respect to the other. For example, in one embodiment, ramps 168A, 170A may not be provided, and endplate 112 may be attached to frame 152.

FIG. 11 illustrates an implant 100 of the disclosure implanted between adjacent vertebrae 10, 12. Frame 152 defines a distal or leading end 152A which is inserted first into the body, and a proximal or trailing end 152B which passes last into the body, the distal and proximal ends defining a longitudinal axis extending therebetween. Implant 100 can be inserted into the body, and into a position between vertebrae, using minimally invasive methods, for example using a small incision, and implant 100 may be passed through a cannula or other structure which maintains a pathway through body tissue. Implant 100 may be inserted into the spinal column through any approach, including anterior, anterolateral, lateral, or posterolateral. A portion of the disc annulus, and nucleus pulposus may be removed in order to form a space into which implant 100 may be inserted. When implant 100 is in a compressed, or reduced height configuration, dovetail guides 200, 202 can be provided to foster maintenance of a relative orientation of upper and lower endplates during insertion or removal of device 100. Dovetail guides 200, 202 further stabilize endplates 110, 112 during expansion, and when implant 100 is expanded. Dovetail guides 200, 202, can have the form of a tongue and groove configuration, or other sliding mating configuration, with ends of ramps 168, 168A, for example.

Implant 100 can be inserted configured to have a lower height profile, as shown in FIG. 2, whereby an extent of distraction of body tissue may be reduced during insertion. Moreover, to the extent that implant 100 is used to open a pathway towards an implantation site, trauma to adjacent tissue is reduced relative to inserting an implant having a final height profile. Once implant 100 is positioned between adjacent vertebrae, actuator screw is rotated by a tool. The tool may be positioned entirely within the body, or can extend from in interior of the body to outside the body, for example having a driving tip at one end and having a handle at an opposite end, with a shaft extending into the body between each end.

Once actuator screw 154 has been rotated to separate endplates 110, 112 a desired amount, the tool is removed. At this point, actuator screw 154 may be secured in place, for example using a mechanical block, or an adhesive, to prevent unintended rotation of actuator screw 154. As carriage 156 is slideably moved by rotation of actuator screw 154, a ramp 166, 166A or a ramped surface of channel 164, 164A of at least one of endplate 110, 112 slides against at least one ramp 168, 168A, 170, or 170A of carriage 156, to cause the endplate to move along an axis transverse to the longitudinal axis of the frame, to increase a height of the implant. Rotation of actuator screw 154 in an opposite direction causes movement along an axis transverse to the longitudinal axis of the frame to decrease a height of the implant.

Polymeric insets, or a polymeric square nut, for example PEEK, can be provided, engageable with threads 158 or other portion of actuator screw 154, to provide additional friction to prevent height loss under load, particularly under cyclic loading. Similarly, once bone screws 300 have been inserted, blocking elements 120 may be rotated to extend over an end of bone screw head 302, preventing screw 300 from backing out. A similar mechanical block (not shown) may be provided for actuator screw 154.

With reference to FIGS. 1-3, 5-8, it may be seen that a socket 118 for a polyaxial screw head 302 can be formed entirely within one of upper or lower endplate 110, 112, or may be formed partially within each of endplate 110 and 112, whereby when implant 100 has been expanded to a final height, the proportions of an interior of socket 118 are

correct or substantially correct for retaining screw head **302**. For example, in FIG. 8, metallic portion **124** forms an upper portion **190** of socket **118**, and mating metallic portion **124A** forms a lower portion **192** of socket **118**. In the embodiment illustrated in the figures, there are three sockets **118**, and all are formed of upper and lower portions. However, there may be more or fewer sockets **118**, and one or more sockets may be formed entirely in an upper or lower endplate.

In an embodiment, implant **100** of the disclosure provides an actuator that translates relative to the body by means of a threaded actuator screw **154**. Ramps **168**, **168A** and **170**, **170A** on a carrier **152** mate with channels **164**, **164A**, and or ramps **166**, on endplates **110**, **112**. Linear translation of carriage **156** causes endplates **110**, **112** to expand implant **100** along an S/I axis with respect to the body. There can be dovetail guides that capture endplates **110**, **112** when collapsing the implant.

Assembly screws **162** fasten endplates made of dissimilar materials, for example PEEK polymeric portions **122**, **122A** to Titanium metallic portions **124**, **124A**. A dovetail and press fit design can be used to connect the dissimilar endplate portions. A PEEK bushing or washer **184** is used between the threaded actuator screw **154** and frame **152** to minimize friction during expansion of implant **100**. Support screws **174** and channels **176** cooperate to form side or lateral stabilizers, and set screw **178** supports a nose or leading end of carriage **156**. Additionally, cooperating slots and projections (not shown) in carriage **156** and frame **152** can be provided for further relative guidance and stability.

In one embodiment, three bone screws **300** are used to provide fixation into adjacent vertebral bodies, two screws **300** passing through implant **100** and into one vertebra, and one screw **300** passing through implant **100** into another vertebra, although other combinations may be used. Bone screws **300** can have spherical or otherwise curved heads, facilitating insertion at a desired angle, or may be provided to mate with socket **118** in a fixed orientation, particularly depending on a diameter of a neck portion of screw **300**. Cam style blocking fasteners **120** can be used to block bone screws **300** from backing out after being inserted.

Implants of the disclosure enable a continuous expansion and retraction over a range of displacements according to predetermined dimensions of a specific implant **100** design. This provides the ability to distract vertebral bodies to a desired height, but also to collapse the implant **100** for repositioning, if therapeutically advantageous for the patient. Endplates **110**, **112** may be shaped to form planes or surfaces which converge relative to each, to provide for lordosis, and can be provided with openings, forming a graft chamber **204** through the openings and between the respective openings through which bone may grow, and into which bone graft material may be placed. Implant **100** may be used to distract, or force bones of a joint apart, or may be used to maintain a separation of bones created by other means, for example a retractor.

Implant **100** may be fabricated using any biocompatible materials known to one skilled in the art, having sufficient strength, flexibility, resiliency, and durability for the patient, and for the term during which the device is to be implanted. Examples include but are not limited to metal, such as, for example titanium and chromium alloys; polymers, including for example, PEEK or high molecular weight polyethylene (HMWPE); and ceramics. There are many other biocompatible materials which may be used, including other plastics and metals, as well as fabrication using living or preserved tissue, including autograft, allograft, and xenograft material.

Portions or all of the implant may be radiopaque or radiolucent, or materials having such properties may be added or incorporated into the implant to improve imaging of the device during and after implantation.

For example, metallic portions **124**, **124A** of endplates **110**, **112** may be manufactured from Titanium, or a cobalt-chrome-molybdenum alloy, Co—Cr—Mo, for example as specified in ASTM F1537 (and ISO 5832-12). The smooth surfaces may be plasma sprayed with commercially pure titanium, as specified in ASTM F1580, F1978, F1147 and C-633 (and ISO 5832-2). Polymeric portions **122**, **122A** may be manufactured from ultra-high molecular weight polyethylene, UHMWPE, for example as specified in ASTM F648 (and ISO 5834-2). In one embodiment, PEEK-OPTIMA (a trademark of Invivo Ltd Corp, United Kingdom) may be used for one or more components of implant **100**. For example, polymeric portions **122**, **122A** can be formed with PEEK-OPTIMA, which is radiolucent, whereby bony ingrowth may be observed. Other polymeric materials with suitable flexibility, durability, and biocompatibility may also be used.

In accordance with the invention, implants of various sizes may be provided to best fit the anatomy of the patient. Components of matching or divergent sizes may be assembled during the implantation procedure by a medical practitioner as best meets the therapeutic needs of the patient, the assembly inserted within the body using an insertion tool. Implants of the invention may also be provided with an overall angular geometry, for example an angular mating disposition of endplates **110**, **112**, to provide for a natural lordosis, or a corrective lordosis, for example of from 0° to 6° for a cervical application, although much different values may be advantageous for other joints. Lordotic angles may also be formed by shaping one or both of plates **110**, **112** to have relatively non-coplanar surfaces. Expanded implant heights, for use in the cervical vertebrae for example, may typically range from 7 mm to 12 mm, but may be larger or smaller, including as small as 5 mm, and as large as 16 mm, although the size is dependent on the patient, and the joint into which an implant of the invention is to be implanted. Implants **100** may be implanted within any level of the spine, and may also be implanted in other joints of the body, including joints of the hand, wrist, elbow, shoulder, hip, knee, ankle, or foot.

In accordance with the invention, a single implant **100** may be used, to provide stabilization for a weakened joint or joint portion. Alternatively, two, three, or more implants **100** may be used, at a single joint level, or in multiple joints. Moreover, implants **100** may be combined with other stabilizing means.

Additionally, implant **100** may be fabricated using material that biodegrades in the body during a therapeutically advantageous time interval, for example after sufficient bone ingrowth has taken place. Further, implant **100** is advantageously provided with smooth and or rounded exterior surfaces, which reduce a potential for deleterious mechanical effects on neighboring tissues.

Any surface or component of the invention may be coated with or impregnated with therapeutic agents, including bone growth, healing, antimicrobial, or drug materials, which may be released at a therapeutic rate, using methods known to those skilled in the art.

Devices of the disclosure provide for adjacent vertebrae to be supported during flexion/extension, lateral bending, and axial rotation. In one embodiment, implant **100** is indicated for spinal arthroplasty in treating skeletally mature patients with degenerative disc disease, primary or recurrent disc

herniation, spinal stenosis, or spondylosis in the lumbosacral spine (LI-SI). Degenerative disc disease is advantageously defined as discogenic back pain with degeneration of the disc confirmed by patient history and radiographic studies, with or without leg (radicular) pain. Patients are advantageously treated, for example, who may have spondylolisthesis up to Grade 1 at the involved level. The surgery position implant 100 may be performed through an Anterior, Anterolateral, Posterolateral, and/or Lateral approach.

In a typical embodiment, implant 100 has a uncompressed height, before insertion, of 12 to 18 mm, and may advantageously be provided in cross-sections of 23×32 mm, 26×38 mm and 26×42 mm, with 4, 8, 12, or 16 degree lordotic angles, although these are only representative sizes, and substantially smaller or larger sizes can be therapeutically beneficial. In one embodiment an implant 100 in accordance with the instant disclosure is sized to be inserted using an MIS approach (a reduced incision size, with fewer and shorter cuts through body tissue).

Implant 100 may advantageously be used in combination with other known or hereinafter developed forms of stabilization or fixation, including for example rods and plates.

Referring now to FIGS. 13-18, implant 100 can be inserted into the intervertebral disc space at a collapsed height, and then expand it to restore the disc space height. Implant 100 provides distraction as well as achieves optimal sagittal balance. As discussed, there are multiple methods and approaches by which implant 100 can be inserted. FIGS. 14-18 illustrate one possible method and approach of the disclosure. While a series of numbered steps are described, it should be understood that there can be numerous other steps pertaining to the procedure, and that the steps described emphasize useful steps in the deployment of implant 100 of the disclosure.

Step 1: Approach—An approach to the desired section of the spine is performed using surgical instruments such as scalpels and retractors, for example using minimally invasive techniques.

Step 2: Preparation—Disc preparation instruments can be used to expose the disc and remove disc material, for example using rongeurs and other suitable instruments (not shown), to create a disc space 14.

Step 3: Trialing—As may be seen in FIG. 14, trialing for implant footprint, height and wedge angle is performed to indicate which size or type of implant 100 is to be used. An expandable trial, static trials, or a combination of each may be used. In FIG. 14, trial implant 320 is trial fit using trial insertion tool 400.

Step 4: Insertion—Graft material or other therapeutically beneficial material is packed into graft chamber 204 of the selected implant 100 when it is collapsed or partially expanded. As may be seen in FIG. 15, implant 100 is inserted into disc space 14 using insertion tool 410. Tool engagement formations 206 are provided on opposite sides of frame 152 or one of endplate 124 or 124A, as can be seen in FIG. 1. Tool arms 412 securely and releasably engage tool engagement formations 206, and align an expansion driver 414 with actuator screw 154.

Step 5: Expansion—In FIG. 16, implant 100 is expanded, as described herein, by turning actuator screw 154 using expansion driver 414. After expansion, additional bone graft material can be packed through graft portals 208 into the central graft chamber 204 using a bone funnel 440 (FIG. 40). A push rod (not shown) can be used for driving graft material through funnel 440.

Step 6: Hole Preparation—Bone screw pilot holes can be formed into one or more adjacent vertebrae, prepared using,

for example, awls, drills and or taps. Multiple pilot holes can be prepared first, or pilot holes can be prepared one at a time, before the insertion of each screw 300. During any of the steps herein, imaging can be carried out to avoid damage to adjacent tissue.

Step 7: Screw Insertion—In FIG. 17, bone screws 300 are inserted using bone screw driver 416. To facilitate access for bone screw driver 416, expansion driver 414 may be withdrawn from insertion tool 410. After bone screws 300 are inserted, they can be blocked from backing out using blocking element 120. Lagging of the vertebral bodies can be performed before or after the bone screws are locked. Fluoroscopy or other imaging can be used to confirm final placement. Imaging can also be used at any of the steps to confirm work performed. Further, bone screw hole preparation and bone screw 300 insertion can be carried out prior to implant 100 expansion, to promote anchoring of the implant during expansion. In FIG. 18, an expanded implant 100 can be seen between vertebrae, secured by bone screws 300. The foregoing method provides a customized fit using implant 100, and minimizes disruption to patient anatomy.

Referring now to FIGS. 19-32, an alternative implant 100B of the disclosure has a shorter actuator screw 154B relative to actuator screw 154 of implant 100 of FIG. 1. Actuator screw 154B engages a proximal end of carriage 156B, and does not pass through graft portal 208B. A compact actuator frame 212 includes a screw bearing 210, and upper and lower tabs 214, 216, respectively. Endplate slots 218, 220 within endplates 110B and 112B slidably receive upper and lower tabs 214, 216. In this manner, actuator screw 154B is rotatably fixed along a longitudinal axis with respect to endplates 110B and 112B, the longitudinal axis indicated in FIG. 19 to extend between distal (“D”) and proximal (“P”) ends. Endplates 110B, 112B can slide upon collar tabs 214, 216 to mutually separate to form an expanded configuration of implant 100B. Actuator screw 154B can be rotatably retained within compact actuator frame 212, so that carriage 156B can be pushed or pulled in threaded engagement with actuator screw 154B, without an axial displacement of actuator screw 154B. This can be accomplished, for example, by a clip or other cooperative engagement between compact actuator frame 212 or bearing 210, and actuator screw 154B, or a blocking element (not shown) partially covering an end portion of actuator screw 154B. In an embodiment, tabs 214 and 216 form a dovetail connection with endplate slots 218, 220.

It should be understood that implant 100 may be identified with a suffix herein, for example 100B, 100C, 100D, 100E, to indicate embodiments illustrating various features of the disclosure. In consideration of the impracticality of illustrating and describing every possible permutation of features, it should be understood that, where logical, features of the various implants may be substituted among the implants. Thus, all of the implants may collectively be referred to as implant 100, unless a specific reference is made to a feature illustrated by a particular embodiment.

Actuator screw 1546B threadably engages carriage 156B at threads 160B, whereby rotation of screw 154B causes carriage 156B to move towards or away from compact actuator frame 212. Carriage 156B has ramps 168, 168A and 170, 170A, which engage corresponding endplate ramps 164, 164A, 166, 166A as described with respect to implant 100. As actuator screw 154B is rotated, carriage 156 translates with respect to endplates 110B, 112B. As a result, carriage ramps 168, 168A and 170, 170A slide against endplate ramps 164, 164A, 166, 166A, causing endplates 110B, 112B to mutually separate. In an embodiment, car-

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riage **156B** is polymeric at threads **160B**, and an interference fit is formed between actuator screw **154B** and threads **160B**, whereby sufficient friction is created to resist unintended rotation of actuator screw **154B**, with a consequential change in height of implant **100B**.

Frame **152** slidably bears against frame support edges **224** extending along endplates **110B**, **112B**, and is slidably connected to carriage **156B** by carriage support screws **174**. In this manner, carriage **156B** is laterally supported, and inhibited from rotational movement, but may move longitudinally along a path defined by carriage support channel **176** and actuator screw **154B**. Additionally, channels or dovetail guides **200**, **202** in endplates **110B**, **112B** receive mating end portions **200A**, **202A** of carriage ramps **168**, **168A**, **170**, **170A**, to further guide and stabilize endplates **110B**, **112B**.

FIGS. **19-32** further illustrate an alternative blocking element **120B**, which, as with other of the various alternative elements herein, may be combined with other implant embodiments herein. Element **120B** forms an sliding block **226** within a block groove **228**, block **226** and block groove **228** forming a dovetail or other sliding mating engagement, wherein block **226** is confined to movement along a path defined by block groove **228**. Once bone screw head **302** is fully seated within bone screw socket **118**, block **226** may be slid partially out of engagement with block groove **228** to a position over bone screw head **302**, thereby blocking a movement of bone screw **300** out of engagement with body tissue. In the embodiment shown, two blocking elements **120B** are illustrated, wherein a tool having two end portions (not shown) can be inserted adjacent each block **226**, and the tool rotated to move both blocks into a blocking position. Accordingly, blocks **226** together form substantially concentric arcs pivoting about the same or close axes.

Implant **100B** is configured to facilitate the insertion of graft material or other therapeutic material through one or more of bone screw socket **118** into graft chamber **204** formed by openings within endplates **110B**, **112B**, and carriage **156B**. After the material is inserted, bone screws **300** may then be inserted into socket **118** and fastened to body tissue as otherwise shown and described herein. A bone funnel **440** (FIG. **40**) may be used to urge material into graft chamber **204**. Alternatively, once implant **100B** is expanded, materials may be inserted into an endplate gap **230** formed by a separation of endplates **110B**, **112B**, as may best be seen in FIGS. **30** and **32**, which are cross-sections taken through compact actuator frame **212**, and upper and lower tabs **214**, **216**.

It should be understood that endplates of the disclosure, in all embodiments, may be formed of a unitary material, as illustrated in FIGS. **19-32** for example, or multiple materials, as illustrated in FIGS. **1-6** for example. Accordingly, endplates **110B**, **112B** may be formed of multiple materials, for example titanium for a proximal, bone screw engaging portion, and UHMWPE for a distal, bone engaging portion. Further, endplates **110B**, **112B** may be provided with teeth or other projections, to positively engage body tissue and reduce a likelihood of undesired migration of implant **100B**.

With reference to FIGS. **33-40**, a spacer implant **100C** includes frame **152C** which forms a dovetail engagement with upper and lower endplates **110C**, **112C**. In this manner, endplates **110C**, **112C** are further stabilized throughout a range of expansion of implant **100C**. As may be seen in FIG. **36**, a cross section of endplate portion **124C** illustrates frame support channel **232** of endplate portion **124C** is shaped to slidably retain frame extension guide **234** of frame **152C** (also visible in FIGS. **44-45**). It should be understood that an

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inverse configuration can be created, wherein a channel is formed in frame **152C** and an extension is formed from endplate portion **124C**. Similar channels and extensions can be formed on opposing sides of frame **152C**, as illustrated, with a frame support channel **232** formed in lower endplate portion **124C'**, as well. In an embodiment, frame **152C** can form an extended region **238** along all or part of the dovetail engagement area of frame support channel **232** and extension guide **234**. For example, frame **152C** can extend in superior and inferior directions to extend from near an outer surface of endplate **110C** to near an outer surface of endplate **112C**, or may extend over a lesser distance. Channel **232** and extension guide **234** are illustrated as transverse to an A-P or longitudinal axis of implant **100C**. In an alternative embodiment, channel **232** and guide **234** are disposed at a non-transverse angle with respect to the longitudinal axis.

With reference to FIGS. **35** and **37**, carriage **156C** includes a graft chamber portal **236**, providing access from a exterior to a proximal end of implant **100C** into graft chamber **204**, after implant **100C** is implanted within the body. Carriage **156C** includes two portals **236** specifically formed to admit the passage of graft or other therapeutic materials, however one or more than two portals **236** can be provided. In the embodiment illustrated, graft chamber portals **236** are formed within a portion of carriage ramp **170**, although other portions of carriage **156C** may be shaped or opened in a like manner. A bone funnel **440** may be used to direct material through one or more of graft chamber portal **236**.

As can be seen in FIG. **35**, actuator screw **154C** includes actuator screw bearing **184C** and lateral screw bearings **240**, provided to promote smooth rotation of actuator screw **154C**. Bearing channels **242** within actuator screw **154C** can be provided to maintain an orientation of lateral screw bearings **240** within screw guide **246** of carriage **156C**. In an embodiment, an interference fit is formed between lateral screw bearings **240** and screw guide **246**, to prevent unintended rotation of actuator screw **154C**. To further stabilize carriage throughout at least a portion of its range of motion, stabilizing posts, screws, or pins **248** can be provided, connected to frame **152C**, for example within frame pin bore **250** by threads, adhesive, or an interference fit, and slideably engageable within pin bores **252** within carriage **156C**. Alternatively, pins **248** can be affixed to carriage **156C**, and can slide within frame pin bores **152C**. In an embodiment,

As can be seen in FIGS. **35** and **38-39**, one or more radiographic markers **254** are positioned within implant **100C**, for example within radiotransparent portions of implant **100C**, or any other radiotransparent portion of the various embodiments herein. For example, a radiographic marker can be positioned within polymeric endplate portion **122**, **122A**, so that an expanded or contracted position thereof may be positively ascertained using imaging. As may be seen in FIG. **39**, radiographic markers **254A**, **254B** are oriented to be aligned with an end of carriage ramps **168**, **168A**, which in this embodiment are radiopaque, only when implant **100C** is fully expanded. To indicate an extent of expansion, one or more radiopaque markers **254** can be positioned with respect to frame **152**, carriage **156**, or any other portion of implant **100** which does not move together with an endplate **110**, **112**, and which is radiopaque, or which is similarly configured with a radiopaque marker **254**.

FIG. **40** illustrates a bone funnel **440** useable with implants **100**, **100B**, **100C**, **100D**, **100E** (collectively, herein, **100**) of the invention. An output aperture **442** is placed proximate an opening into an open area within implant **100**, for example graft chamber **204**. Bone graft material, and or

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other therapeutic agents, are collected within including for example bone growth factors, antimicrobial agents, or other therapeutic is placed into widened input chamber 444, and then pushed down pipe 446 with a driver, for example a rod (not shown). A pipe connector 448 can be provided, sized to correspond to graft chamber portal 236. Driven bone graft material is passed into an interior of implant 100, where it may have its intended therapeutic benefit upon contacting body tissue of at least one vertebra.

In an embodiment, carriage ramps 168, 168A, 170, 170A can have differing ramp angles and or sizes, wherein endplate ramps 166, 166A have corresponding profiles and sizes. For example, if ramps 168, 168A are shorter than ramps 170, 170A, expansion will occur at a greater rate along a proximal side of implant 100, and in this manner an angular orientation of the spine, for example lordosis, may be corrected. Similarly, ramps 170, 170A can be shorter than ramps 168, 168A. Alternatively, one side of ramp 168, 168A can be shorter than another side of ramp 168, 168A, with a corresponding difference along ramps 170, 170A. In this manner, a sideways orientation of the spine, for example Scoliosis, may be corrected.

FIGS. 41-43 illustrate an alternative implant 100D of the disclosure, which pivots proximate ends of endplates 110D, 112D, providing both axial translation, as indicated by arrows "A", and pivoting, as indicated by arrows "B". Axial translation is maintained using frame 152C, together with frame extension guide 234 and frame support channel 232, as described with respect to implant 100C. However, an endplate pivot 256 is formed between endplate portions 122D and 124D, and between endplate portions 122D' and 124D'. FIG. 43 illustrates implant 100D with frame 152C removed, illustrating an endplate hinge 258 formed between endplate portions 122D and 122D'. Connected in this manner, endplate portions 122D and 122D' pivot about endplate hinge 258, as well as endplate pivots 256. Accordingly, a height of implant 100D at a distal end of implant portions 122D and 122D' is held constant, while a proximate end of implant portions 122D and 122D' translates axially with endplate portions 124D and 124D' to increase a height of implant 100D.

Implant 100D can be inserted into the intervertebral disc space at a collapsed height, and then expanded into lordosis to restore sagittal balance and height loss in the disc space. Implant 100D provides distraction as well as achieving optimal sagittal balance. Further, implant 100D reduces impaction to body tissue during insertion at a collapsed height, and gives a medical practitioner the capability to continuously adjust the lordotic angle of the supporting endplates to best fit the patient's anatomy and therapeutic needs.

Endplate pivot 256 is formed as mating circular portions of endplate portions 122D and 124D, and of endplate portions 122D' and 124D'. While one endplate portions forms an extension, and the other a receptacle, it should be understood that this configuration may be reversed.

Endplate hinge 258 is formed as a flexible connector 260 extending between endplate portions 122D and 122D'. In an embodiment, endplate portions 122D and 122D' are molded as a single part from a polymeric or other flexible material, thus forming a living hinge. In a further embodiment, a hinge is formed between endplate portions 122D and 122D' by any known means, including a barrel or flag hinge, or a hinge similar in style to endplate pivots 256. In an alternative embodiment, endplate hinge 258 is formed in connection with frame 152C.

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By providing both axial and pivoting movement of endplate portions, implant 100D enables the formation of an alternative supporting structure, and in particular, a supporting structure with a convex conformity. This can be useful to correct particular spinal problems, including lordosis, for example.

With reference to FIGS. 44-45, which are cross-sections of an alternative implant 100E of the disclosure, it may be seen that actuator screw 154E is rotatably connected to frame 152E, for example using C-clip 262, as illustrated. An alternative method of rotatably securing actuator screw to frame 152E can include, for example, a leading set screw 178 (see, e.g. FIGS. 6, 6A) that freely spins relative to frame 152E, but is affixed to actuator screw 154E. An alternative method includes forming mating portions (not shown) upon frame 152E and screw 154E.

Further stability can be provided for carriage 156C through the use of stabilizing pins 248, frame pin bores 250, and pin bores in carriage 152C, as described with respect to implant 100C herein.

In a further embodiment, actuator screw 154E' is shorter than actuator screw 154E, and thereby reduces an obstruction of graft chamber 204. A tool can be passed through screw guide 246, and then through graft chamber 204, to engage actuator screw proximal end 182. Graft material can additionally be passed through screw guide 246, and placed within graft chamber 204. Bone funnel 140 can be used to pass materials through screw guide 246, and pipe connector can be adapted or replaced to best fit the dimensions of screw guide 246.

FIG. 46 depicts an alternative expandable implant including guide pins in accordance with embodiments of the present application. The expandable implant 100 includes many features disclosed in prior embodiments, including a first endplate 110, a second endplate 112, a frame 152 for receiving an actuator 150 therein, and an actuator screw 154. In addition, the expandable implant 100 in FIG. 46 includes additional features, including guide pins 359, first and second compressible clips 380, 382 and a blocking mechanism 382.

As shown in FIG. 46, the expandable implant includes an upper or first endplate 110 and a lower or second endplate 112. The upper endplate 110 can be comprised of a first portion 122A and a second portion 124A that are operatively coupled together (e.g., via a fastener). In some embodiments, the first portion 122A comprises a polymeric portion, while the second portion 124A comprises a metallic portion. By having an endplate formed of a polymeric portion and a metallic portion, this advantageously provides an endplate that is both radiolucent and strong. The lower endplate 112 can be comprised of a first portion 122B and a second portion 124B that are operatively coupled together (e.g., via a fastener). In some embodiments, the first portion 122B comprises a polymeric portion, while the second portion 124B comprises a metallic portion. In other embodiments, the endplates 110, 112 can be formed of a single piece that is formed of either a polymer, such as PEEK, or a metal, such as titanium or cobalt-chrome. As in prior embodiments, the upper endplate 110 and the lower endplate 112 can include one or more bore holes for receiving bone screws therein. For example, as shown in FIG. 46, lower endplate 112 includes at least one bore hole 189 for receiving a bone screw therethrough for securing the implant 100 to an adjacent bone member.

A frame 152 for receiving an actuator 150 is positioned between the first endplate 110 and the second endplate 112. The frame 152 is configured to receive side support screws

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174 through channels to secure the frame 152 to the actuator 150. In addition, one or more guide pins 359 are provided at a distal or leading end of the frame 152. The guide pins 359 are inserted through openings in the frame 152 and contact a surface of the actuator 150. By engaging the actuator 150, the guide pins 359 advantageously stabilize the actuator 150 such that it is not tilted during use.

The actuator 150 comprises a moveable carriage 156 having a first pair of upper ramped surfaces 170 connected to a second pair of upper ramped surfaces 168 via a bridge member 199. In some embodiments, the first pair of upper ramped surfaces 170 and the second pair of upper ramped surfaces 168 are inclined in the same direction (e.g., toward the distal or leading end of the actuator 150). The first and second pair of upper ramped surfaces 170, 168 are configured to engage corresponding angled or ramped surfaces of the first endplate 110, such that movement of the carriage 156 causes expansion of the implant 100. In some embodiments, a first pair of lower ramped surfaces extends downwardly from the first pair of upper ramped surfaces 170, while a second pair of lower ramped surfaces extends downwardly from the second pair of upper ramped surfaces. The first and second pair of lower ramped surfaces are configured engage corresponding angled or ramped surfaces of the second endplate 112, such that movement of the carriage 156 causes expansion of the implant 100.

An actuator screw 154 can be provided to actuate the actuator 150. The actuator screw 154 comprises a head portion 197 and a shaft portion 198. The shaft portion 198 comprises threads for engaging a corresponding threaded portion of the actuator 150. Rotational movement of the actuator screw 154 in a first direction causes linear translation of the moveable carriage 156 of the actuator 150, thereby causing separation of the endplates 110, 112 and expansion of the implant. Rotational movement of the actuator screw 154 in a second direction opposite the first direction causes linear translation of the moveable carriage 156 of the actuator 150 in an opposite direction, thereby causing contraction of the endplates 110, 112.

To retain the actuator screw 154 in the implant 100, an actuator frame 212 can be provided. The actuator frame comprises an upper tab portion 214 and a lower tab portion 216, and an opening 213 therebetween for receiving the actuator 150 therethrough. The upper tab portion 214 can be received in a slot in the first endplate 110, while the lower tab portion 216 can be received in a slot in the second endplate 112. To maintain the actuator 150 in the actuator frame 212, a first compression clip or "C-clip" locking mechanism 380 can be provided to secure the actuator 150 to the actuator frame 212. The C-clip 380 is configured to fit around a portion of the actuator 150, such as the head portion 197. In some embodiments, the C-clip 380 is retained within a recess or groove formed around the circumference of the actuator head portion. The C-clip 380 is advantageously configured to compress such that it can be trapped in a recess 288 in the actuator frame 212 (as shown in FIGS. 47A and 47B). This advantageously secures the actuator 154 within the actuator frame 212.

To prevent the bone screws from inadvertently backing out, a blocking mechanism 390 can be provided and attached via a C-clip 382. The blocking mechanism 390 comprises a body 391 having an opening 392 for receiving at least a portion of the head portion 197 of the actuator screw 154 therethrough. In some embodiments, a second compression clip or "C-clip" locking mechanism 382 can be provided to secure the blocking mechanism 390 to the actuator screw 154. As shown in FIGS. 47A and 47B, the C-clip 382 is

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configured to fit around the head portion 197 of the actuator screw 154. In some embodiments, the C-clip 382 can be received within a recess or groove formed in the head portion 197 of the actuator screw 154. The C-clip 382 is configured to compress until it reaches a recess 289 (shown in FIGS. 47A and 47B) in the blocking mechanism 390, thereby securing the C-clip 382 to the blocking mechanism 390.

FIGS. 47A and 47B show the implant 100 of FIG. 46 in unexpanded and expanded configurations, respectively. From these views, one can see the additional novel features, such as the first C-clip 380 and the second C-clip 382 retained in recesses 288, 289, thereby maintaining the components of the implant 100 in a secure connection.

In some embodiments, the implant 100 of FIG. 46 is sized and configured to be used in an anterior approach. In some embodiments, when assembled, the leading end of the implant 100 comprises a convex surface, while the trailing end of the implant 100 comprises a substantially flat surface. The leading end and the trailing end can be separated by curved arms, formed by side arms of the frame 152.

A method of insertion is now provided. After forming an incision in a patient and removing tissue from a disc space, a surgeon can insert the implant 100 through an anterior approach. The implant 100 can be inserted in an unexpanded configuration, as shown in FIG. 47A. Once the implant 100 has been inserted into the disc space, the implant 100 can be expanded by rotating or actuating the actuator screw 154 (e.g., via a driver). This causes translational movement of the carriage 156 with the ramps, thereby causing expansion of the implant 100. In some embodiments, prior to inserting and expanding the implant 100, bone graft material can be provided in a graft opening of the implant. In other embodiments, the implant 100 can include an opening that can receive bone graft therethrough after inserting the implant 100 in a disc space. In some embodiments, bone graft material can be inserted through the screw opening 189 (shown in FIG. 46) after the implant 100 is inserted into a disc space. In other embodiments, the implant 100 can be used in different approaches, including posteriorly or laterally.

FIG. 48 depicts an alternative expandable implant including endplates that are formed primarily of metal in accordance with embodiments of the present application. The expandable implant 100 includes many features disclosed in prior embodiments, including a first endplate 110, a second endplate 112, a frame 152 for receiving an actuator 150 therein, and an actuator screw 154. The expandable implant also includes guide pins 359 and compression C-clips 380, 382 as discussed with respect to prior embodiments.

As shown in FIG. 48, the first endplate 110 is a single-piece member formed of a metal. Likewise, the second endplate 112 is a single-piece member formed of a metal. In some embodiments, the first and second endplates 110, 112 are formed completely of a metal, as shown in FIG. 48. In other embodiments, the first and second endplates 110, 112 are formed primarily of a metal, but may have traces of other materials. Advantageously, by providing endplates that are formed primarily or completely of metal, this advantageously creates an implant having increased strength.

In some embodiments, the expandable implant 100 in FIG. 48 is sized and configured to be implanted anteriorly. In other embodiments, the expandable implant 100 can be inserted posteriorly or laterally.

FIGS. 50A and 50B illustrate unexpanded and expanded configurations of an alternative expandable implant having an alternative means to capture the blocking mechanism in

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accordance with embodiments of the present application. The expandable implant 100 includes many similar features as prior embodiments, including a first endplate 110, a second endplate 112, a frame 152 for receiving an actuator 150, and an actuator screw 154. The implant 100 in the current embodiment, however, has a distinct blocking mechanism 390 that is not attached to the actuator screw 154 by a C-ring. Rather, the blocking mechanism 390 is captured between the actuator screw 154 and the actuator plate 212. The blocking mechanism 390 includes an extension portion 394 that can fit in a recess, groove or track formed in the actuator plate 212. As the actuator screw 154 is rotated and linearly translated, the actuator screw 154 pushed in the blocking mechanism 390 such that the extension portion 394 of the blocking mechanism 390 engages the track of the actuator plate 212, thereby securely capturing the blocking mechanism 390 within the assembly. In other embodiments, a compression C-ring can be optionally provided to retain the blocking mechanism 390 on the actuator screw 154, in addition to the engagement mechanism discussed herein.

All references cited herein are expressly incorporated by reference in their entirety. There are many different features to the present invention and it is contemplated that these features may be used together or separately. Unless mention was made above to the contrary, it should be noted that all of the accompanying drawings are not to scale. Thus, the invention should not be limited to any particular combination of features or to a particular application of the invention. Further, it should be understood that variations and modifications within the spirit and scope of the invention might occur to those skilled in the art to which the invention pertains. Accordingly, all expedient modifications readily attainable by one versed in the art from the disclosure set forth herein that are within the scope and spirit of the present invention are to be included as further embodiments of the present invention.

What is claimed is:

1. A surgical method comprising:
forming an incision in a patient;
removing tissue from the patient at a surgical site; inserting an expandable implant into the surgical site via an anterior approach; and
expanding the expandable implant at the surgical site wherein the expandable implant comprises an upper endplate, a lower endplate, a moveable actuator positioned therebetween, and a rotatable actuator screw configured to actuate the moveable actuator,
wherein the expandable implant further comprises a first bone fastener insertable through a first through hole in the upper endplate and a second bone fastener insertable through a second through hole in the lower endplate.
2. The surgical method of claim 1, wherein the upper endplate is formed of a polymeric portion and a metallic portion.
3. The surgical method of claim 1, wherein the upper endplate is formed completely of a metal.
4. The surgical method of claim 1, further comprising inserting the first bone fastener through the expandable implant into a first vertebral body and inserting the second bone fastener through the expandable implant into a second vertebral body.
5. The surgical method of claim 1, wherein the expandable implant comprises an upper endplate, a lower endplate, a moveable actuator with ramps positioned therebetween and an actuator screw, wherein rotation of the actuator screw

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causes translation of the moveable actuator, thereby separating the upper endplate from the lower endplate.

6. The surgical method of claim 5, wherein the expandable implant further comprises an actuator plate for receiving the moveable actuator therethrough.

7. The surgical method of claim 6, wherein the actuator plate includes an upper tab for inserting into the upper endplate and a lower tab for inserting into the lower endplate.

8. The surgical method of claim 6, wherein the expandable implant further comprises a compression clip positioned between the actuator screw and the actuator plate.

9. The surgical method of claim 8, wherein the expandable implant further comprises a second compression clip positioned around the actuator screw.

10. A surgical method comprising:

forming an incision in a patient; removing tissue from the patient at a surgical site;

inserting an expandable implant into the surgical site via an anterior approach;

expanding the expandable implant at the surgical site;
inserting a first bone fastener in the expandable implant to fix the expandable implant to a first bone member; and
inserting a second bone fastener in the expandable implant to fix the expandable implant to a second bone member

wherein the expandable implant comprises an upper endplate, a lower endplate, a moveable actuator positioned therebetween, and a rotatable actuator screw configured to actuate the moveable actuator,

wherein the expandable implant further comprises a first through hole in the upper endplate and a second through hole in the lower endplate for receiving the first and second bone fasteners.

11. The surgical method of claim 10, wherein the actuator comprises a first pair of upper ramps connected to a second pair of upper ramps.

12. The surgical method of claim 11, wherein the first pair of upper ramps and the second pair of upper ramps are configured to engage angled surfaces formed in the upper endplate.

13. A surgical method comprising:

forming an incision in a patient; inserting an expandable implant into the surgical site via an anterior approach;
expanding the expandable implant at the surgical site;
inserting a first bone fastener in the expandable implant to fix the expandable implant to a first bone member; and
inserting a second bone fastener in the expandable implant to fix the expandable implant to a second bone member

wherein the expandable implant comprises an upper endplate, a lower endplate, a moveable actuator positioned therebetween, and a rotatable actuator screw configured to actuate the moveable actuator,

wherein the expandable implant further comprises a first through hole in the upper endplate and a second through hole in the lower endplate for receiving the first and second bone fasteners.

14. The surgical method of claim 13, wherein the expandable implant comprises a convex leading end and a flat trailing end.

15. The surgical method of claim 13, wherein the expandable implant comprises an upper endplate formed of at least two materials.

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